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15 TITLE:

USE OF THE CRYSTAL STRUCTURE OF BACTERIAL

IMP DEHYDROGENASE TO DESIGN INHIBITORS OF

**BACTERIAL GROWTH** 

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### USE OF THE CRYSTAL STRUCTURE OF BACTERIAL IMP DEHYDROGENASE TO DESIGN INHIBITORS OF BACTERIAL GROWTH

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#### 10 BACKGROUND OF THE INVENTION

The invention relates the crystal structure of IMPDH isolated from bacteria. The structure is different from the structure of mammalian or fungal IMPDH, allowing design of selective inhibitors of bacterial IMPDH.

Inosine monophosphate dehydrogenase (IMPDH; Enzyme Comission (EC) 1.1.1.205) is a rate-limiting enzyme in the synthesis of guanine ribonucleotides. IMPDH has an essential role in providing critical precursors for DNA and RNA biosynthesis and in signal transduction pathways that mediate cell differentiation (Collart *et al.*, 1990; Kiguchi *et al.*, 1990). Because of its central role in purine metabolism, IMPDH is an attractive therapeutic target. Several recent reviews have outlined the utility of mammalian IMPDH inhibitors as anticancer (Pankiewicz, 1997) or antiviral (Andrei *et al.*, 1993) agents or as immunosuppressive drugs (Halloran, 1996) (see Table 1).

Table 1: Clinically Useful Inhibitors of IMPDH

	Inhibitor	Clinical Application				
30	Ribavirin	Antiviral				
30	Mycophenolate mofetil	Immunosuppression				
	Mizoribine	Imunosuppression				
35	Tiazofurin	Anticancer				

Although there are no reports of selective inhibitors of bacterial IMPDH enzymes, such compounds could have potential application as specific antimicrobial agents.

The active form of IMPDH enzymes (50-55 kDa) is a homotetramer with four active sites per tetramer. A cysteine residue in the active site forms a covalent

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intermediate with IMP (Wang et al., 1996). A consensus sequence of thirteen amino acid residues that includes cysteine in this active site has been proposed as a signature motif (i.e., an amino acid sequence that can be used as a fingerprint or specific identifier for this class of enzymes) for the IMPDH and guanosine monophosphate (GMP) reductase enzymes (Bairoch, 1995). This IMPDH consensus region is highly conserved in both bacteria and eukaryotes, with 90% and 85% of the respective residues being identical within each kingdom. However, only 40% of these residues remain identical when compared between the two kingdoms. This limited conservation suggests that bacterial and eukaryotic IMPDH enzymes may have distinct characteristics; a suggestion supported by their kinetic differences and differential sensitivity to inhibitors. Enzymes from mammalian sources show distinctly lower values for the K<sub>m</sub> for nicotinamide adenine dinucleotide (NAD) than do those enzymes from bacteria. In addition, mammalian IMPDH enzymes are 10-100 times more sensitive to inhibition by mycophenolic acid (MPA) than are bacterial IMPDH enzymes. Sequence analysis of all known IMPDH enzymes supports the distinction between bacterial and eukaryotic enzymes. A deep branching of the bacterial and eukaryotic forms of IMPDH is observed upon phylogenetic analysis of the relationships among the various IMPDH genes (Collart et al., 1996 a and b). This phylogenetic analysis indicates a general functional conservation of amino acid and suggests a unique amino acid sequence signature for these kingdoms.

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The elucidation of a kingdom-specific signature for IMPDH enzymes is an important element in the development of specific inhibitors. The two partial structures of IMPDH from Chinese hamster (Sintchak et al., 1996) (85% structure complete with bound transition state analogue and mycophenolic acid, MPA) and Tritrichomonas foetus (Whitby et al., 1997) (68% structure complete with bound xanthosine monophosphate [XMP]) have been reported with only the coordinates of the latter available in the Protein Data Bank (PDB). These structures furnished the initial information about the structure and reaction mechanism of eukaryotic IMPDH enzymes. Inhibitors of IMPDH in bacteria are needed to treat infections, in particular, to overcome the barrier of antibiotic resistance.

### **BRIEF SUMMARY OF THE INVENTION**

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The invention relates for the first time a crystal structure of a bacterial IMPDH. This invention relates that bacterial and mammalian IMPDH enzymes provide the same catalytic function, but have a set of unique structural and biochemical characteristics. An embodiment is a crystal structure of IMPDH isolated from Streptococcus pyogenes. S. pyogenes IMP dehydrogenase represents the class of bacterial IMPDH enzymes that show distinct functional differences when compared to mammalian IMPDH enzymes. The bacterial enzymes bind NAD poorly (Zhou et al., 1997; Kerr et al., 1997) (K<sub>M</sub> > 1 mM) and are inhibited by MPA only at very high concentrations (Ki>0.5 mM). Elucidation of the structural basis of these distinct characteristics is useful to aid in design of specific IMPDH inhibitors that will inhibit the infectious agent without harming the host's IMPDH.

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The coding sequence of bacterial IMPDH specifies a protein of 493 amino acids that contain only a single cysteine residue at the active site (Ashbaugh et al., 1995). IMPDH from S. pyogenes is a representative bacterial enzyme because the organism is pathogenic, and therefore a good model for the investigation of enzyme inhibitors. Streptococci are the most common cause of worldwide pneumonia and a leading cause of pediatric infections. The structure of the S. pyogenes bacterial IMPDH provides the basis for elucidation of the structural characteristics that distinguish bacterial from eukaryotic IMPDH enzymes. Knowledge of these characteristics permits an understanding of why these enzymes exhibit functionally distinct behavior and therefore provides a foundation for the design of specific inhibitors of IMPDH that have clinical value.

In addition to inhibiting pathogens, the immunosuppressive use of IMPDH inhibitors is applicable to treat chronic inflammatory diseases such as arthritis, diabetes, or systemic lupus erythromotosis. Use of the IMPDH structure from S. pyogenes will facilitate identification of other pathogens that will be inhibited by drugs that inhibit S. pyogenes.

### **Definitions and Abbreviations**

A "binding pocket" is a space in a molecule in which an inhibitor of the molecule is bound. 30

The following abbreviations are used throughout the application:

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Ala Alanine Thr Threonine A = Val C Valine Cys Cysteine V Y Tyr Tyrosine Leu Leucine L N Ile Asparagine Isoleucine Asn Ι **Proline** Q Gln Glutamine 5 P Pro = = = D F Phe Phenyalanine Asp Aspartic Acid = W Trp Trytophan E Glu Glutamic Acid == == Methionine K Lysine Met Lys M == == G Glycine R Gly Arginine Arg == = Ser His Histidine 10 S Serine H =

**CBS** Cystathionine-β-synthase =Guanosine monophosphate **GMP** == **IMP** Inosine monophosphate =Inosine monophosphate dehydrogenase **IMPDH** Mycophenolic acid **MPA** Nicotinamide adenine dinucleotide NAD = Protein Data Bank **PDB** =**XMP** Xanthosine monophosphate

#### **BRIEF DESCRIPTION OF THE DRAWINGS**

FIG. 1 is a ribbon drawing of the catalytically active IMPDH tetramer; the tetramer is displayed parallel (FIG 1a) and perpendicular (FIG 1b) to the four-fold axis. Each subunit is shown with a spacefilling model of IMP, the active site of each subunit.

FIG. 2 is a representation of the secondary structure of the IMPDH monomer. (FIG 2a) Topology diagram of IMPDH domains. Secondary structure was assigned using the Kabsch and Sander (1983) algorithm along with visual inspection. The  $\alpha$  helices and  $\beta$  strands that form the TIM barrel fold are labeled  $\alpha$ 1- $\alpha$ 8 and  $\beta$ 1- $\beta$ 8. The remaining strands and helices are designated in alphanumeric order (e.g.,  $\alpha_A$ - $\alpha_L$ ). The part of the structure not visible in the electron density maps is marked as "????". (FIG 2b) Stereoview ribbon diagram arranged approximately perpendicular to the axis of the TIM barrel fold; IMP is shown as a ball and stick model.

FIG. 3 shows an IMPDH active site. (FIG 3a) is a cartoon of bound IMP showing side chain interactions and active site residues. (FIG 3b) is a stereoview ball and stick diagram of bound IMP illustrating the alignment of the hypoxanthine ring relative to the catalytic Cys310 residue. The dashed ring cartoon indicates the

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proposed realignment of the hypoxanthine ring initiated by NAD binding. Residues targeted for mutagenesis (E421 and Y450') are underlined. The "'" symbol on Y450 indicates a symmetry related molecule.

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FIG. 4 is a stereoview of the electron density map around the active site; the solvent-flattened MAD map is calculated at 2.5 Å resolution drawn at a contour level of  $1.2\sigma$ .

FIG. 5 is a representation of the secondary structure of the CBS dimer domain shown as a stereoview ribbon diagram arranged approximately along the dyad axis of the two CBS motifs.

#### **DETAILED DESCRIPTION OF THE INVENTION** 10

The invention relates the crystal structure of a bacterial IMPDH with substrate bound in the catalytic site. Conditions for producing a crystal from bacterial IMPDH were completely different than for humans. The structure was determined using SeMet-substituted protein and multi-wavelength anomalous diffraction (MAD) (Hendrickson, 1991) analysis of data obtained with synchrotron radiation from the undulator beamline of the Structural Biology Center at the Advanced Photon Source. The high quality of the data allowed determination of the structure of both catalytic and cystathionine- $\beta$ -synthase (CBS) dimer domains. The  $\alpha/\beta$  barrel domain of IMPDH embodies the catalytic framework. The CBS dimer domain contains two CBS motifs that are known to play a regulatory role in other proteins. However, their function in IMPDH is unknown. This is the first crystal structure reported of a complete CBS dimer domain. Bacterial and mammalian IMPDH enzymes have distinct kinetic and biochemical characteristics. Comparison of this bacterial IMPDH with the known partial structures from eukaryotic organisms provides an explanation of their distinct properties and contributes to the design of specific bacterial inhibitors.

#### **Structure of Bacterial IMPDH**

The structure of S. pyogenes IMPDH (FIG. 1) provides a new resource to define the distinct characteristics of bacterial and mammalian IMPDH enzymes. Features such as the catalytic motifs, active site flap region and CBS dimer domain are structurally conserved, but show a different pattern of sequence conservation in bacteria and eukaryotes. Analysis of sequence differences in these regions suggests they could contribute to the differential signature of the bacterial and mammalian

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enzymes. One of these sequence regions is the  $\alpha G$  helix (FIG. 2) that forms part of the catalytic pocket. Analysis of sequence alignments for this region (Table 2) indicates a pattern of catalytic residues conserved in all enzymes and a secondary pattern of amino acid conservation associated with either bacterial or eukaryotic IMPDH enzymes. In this region, the pattern of bacterial sequence conservation is superimposed on a pattern of residues highly conserved in IMPDH enzymes from all organisms. These highly conserved residues are involved in IMP binding; the characteristics of which appear to be similar for bacterial and eukaryotic IMPDH enzymes. The existence of distinct bacterial catalytic pocket is supported by sitespecific mutants at positions E421 and Y450 (Numbering corresponds to the amino acid sequence of the S. pyogenes IMPDH enzyme) that appear to differentially alter the activity of the mammalian and bacterial IMPDH enzymes. Residue Y450 in S. pyogenes IMPDH is located at the noncatalytic end of the TIM barrel. However, this region has contacts with another molecule in the tetramer and contributes to the catalytic environment of the adjacent monomer (FIG. 3). Site-specific mutagenesis results show partial retention of activity with an alanine substitution but no activity with an aspartic acid substitution for this residue. Aspartic acid was selected as a replacement on the basis of sequence alignments that show 12 of 13 eucaryotic enzymes contain aspartic acid at the corresponding position (the exception being asparagine in T. foetus). The partial activity observed with the Ala replacement suggests Y450 does not have an essential role in catalysis but does contribute to the environment of the catalytic pocket. Further analysis of this region will provide insight into the differences in the environment of the catalytic pocket in bacterial and eucaryotic enzymes and also the role of the tetrameric form of the active enzyme.

The E421 in *S. pyogenes* IMPDH is conserved in bacteria while eucaryotic IMPDH enzymes contain glutamine in the corresponding position. In hamster IMPDH, the corresponding residue, Q441, is implicated in the binding of MPA. Comparison of the residues involved in MPA binding in the hamster enzyme (D274, Ser276, N303, R322, G326, T333, Q441) with the equivalent residues in *S. pyogenes* IMPDH indicates that these residues are largely conserved. The aspartic acid, asparagine, glycine, and threonine residues are identical, but threonine replaces S276 (although serine is present in other bacterial enzymes), and K301 replaces the hamster

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R322 residue. The most significant change appears to be replacement of E421 with Q441 (interestingly, this residue is part of the active-site flap). Although this suggests that the NAD binding pockets of hamster and bacterial IMPDH differ, a change in activity was not observed upon substitution of glutamine for glutamic acid at position 421. It is possible that this substitution does not affect the observed activity but may alter the sensitivity to MPA.

The active site flap represents another region that could account for the kinetic and biochemical differences between IMPDH enzymes. This flap is present in all IMPDH enzymes and is disordered in all IMPDH structures but may become ordered upon NAD binding. Sequence comparisons (Table 2) indicate the loop size is conserved but sequence conservation is limited. A conserved feature of this region is the presence of an Arginine next to one or two aromatic residues. Since IMP and NAD bind sequentially to the active site, these residues may bind to the phosphate or the adenine or nicotinamide ring thereby ordering the active site. The sequence heterogeneity observed in this flap region may also account for the discriminatory features of bacterial and mammalian IMPDH enzymes.

The finger region and the CBS dimer domain are not involved in catalysis but are found in all IMPDH enzymes. These regions show little sequence conservation but have been structurally conserved. The finger structure is composed of two antiparallel β-strand structures stabilized by hydrogen bonding and interactions with the βL region (FIG. 2). The CBS dimer domain contains two CBS motifs arranged on a pseudo-dyad axis. In other proteins (e.g. cystathionine-beta-synthase and chloride channel proteins), mutations in these domains are associated with pathologic consequences. It has also been suggested (Nimmesgern *et al.*, 1996) that these domains may be involved in cytoplasmic targeting or other regulatory functions. In either case, the metabolic expenditure required for conservation of these structures suggests an underlying functional role.

A unique aspect of the *S. pyogenes* IMPDH structure is that it allows examination of the initial stage of the catalytic cycle. IMP does not form a covalent bond in the absence of NAD. Covalent bond formation requires reorientation of the hypoxanthine ring and nucleophilic attack on C2 by Cys310. This suggests that NAD may have multiple roles as hydride acceptor, substrate activator, and also in

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contributing to the structure of the active site pocket. NAD binding likely initiates realignment of the hypoxanthine ring and also facilitates the electron shift with the ring required for formation of the thioimidate intermediate.

The structure of *S. pyogenes* IMPDH allows for a detailed comparison of the eukaryotic and bacterial enzymes and provides the basis for an explanation for the unique properties of the bacterial enzymes. This knowledge aids the design of inhibitors that specifically target bacterial IMPDH enzymes.

### **Determination of Bacterial IMPDH Crystal Structure**

The crystal structure of S. pyogenes IMPDH was determined at 1.9 Å resolution by replacing all methionine residues in the enzyme with selenomethionine and applying MAD phasing methods (Hendrickson, 1991). The IMPDH crystals were tetragonal (space group I422, a = b = 151.49 Å, c = 101.67 Å,  $\alpha = \beta = \gamma = 90^{\circ}$ ) and contained one 53 kDa monomer per asymmetric unit. The enzyme contains 13 methionine residues: a potential 13 selenium sites (Table 3). Data were collected at three x-ray energies: at the peak and edge energies of the selenium absorption spectrum, and at a lower energy far from the edge. The initial model of the  $\alpha/\beta$  barrel core (amino acid residues 15-90 and 222-460) was obtained by molecular replacement using a search model derived from the atomic coordinates of IMPDH from T. foetus (Whitby et al., 1997). These phases were sufficient to permit location of 6 selenium sites. Four rounds of phase development, in which the selenium sites were refined against the three data sets in program MLPHARE (Otwinowski, 1991), permitted location of the remaining selenium atoms in different Fourier maps. The electron density map used for interpretation of this structure was phased by MLPHARE with all 13 selenium sites. The figure of merit (FOM) for this phasing calculation was 0.64 (Table 4); the phasing power was 2.1 for all data between 10-2.5 Å resolution. Solvent flattening and density modification (Cowtan, 1994) further improved the electron density map (FOM=0.72), which at this point was clear enough to trace almost all of the main peptide chain and most side chains (FIG. 4). Registration of the sequence was made easy because methionine residues could be identified with the known selenium positions. The model was refined with the program CNS (Brünger, et al., 1998), which significantly improved the interpretation of several regions in the model that were initially ambiguous. The model disclosed herein contains 3,992

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nonhydrogen atoms, from residues 2-401 and 416-492, in two distinct domains. The crystallographic R-factor is 23.2% (R-free: 26.1%) for all reflections between 6 and 1.9 Å resolution (Table 5). The current model contains 422 water molecules with an average B factor for all nonhydrogen protein atoms of 37.5 Å (Table 5). This structure is significantly more complete (97%) and of higher resolution (1.9Å) than those reported for IMPDH from Chinese hamster (Sintchak *et al.*, 1996) (85%, 2.3 Å) and *T. foetus* (Whitby *et al.*, 1997) (68%, 2.3 Å). The map also contains clearly defined electron density for the IMP substrate, bound in the catalytic site.

### Catalytic Domain of Bacterial IMPDH

The *S. pyogenes* IMPDH tetramer is composed of four identical subunits where each monomer has a two-domain structure (FIG. 1a). The catalytic domain (amino acid residues 2-92 and 224-492) forms the interior core of the active tetrameric enzyme and is approximately 40x40x50 Å. This domain contains the catalytic site that is positioned near the tetramer four-fold at the subunit interface (FIG. 1b). This location places access to the active site on the same face of the tetramer. The CBS dimer domain (residues 93-223, approximately 20x20x30 Å) is on the active site face and projects outward from the core of the tetrameric unit placing this domain in the corner of the square formed by E162.

The core of the catalytic domain (FIG. 2a) is formed by an  $\alpha/\beta$  barrel structure that provides a scaffold for the attachment of additional structural and catalytic moieties and the CBS dimer domain. This core region contains a series of eight parallel  $\alpha/\beta$  motifs with the active site near the C-terminus of the  $\beta$ -strands (FIG 2b). The number and relative location of the barrel structures in *S. pyogenes* IMPDH are similar to that reported for the Chinese hamster (Sintchak *et al.*, 1996) and *T. foetus* (Whitby *et al.*, 1997) IMPDH and for other nicotinamide-dependent oxidoreductases. However, in IMPDH the phosphate-binding site is occupied by IMP rather than by the phosphate of the NAD or NADP cofactors as seen in the other nicotinamide dependent oxidoreductases.

The  $\beta$ -strand structures and the interior residues of the helices are hydrophobic with very few water molecules observed in the interior of the  $\alpha/\beta$  barrel structure. This hydrophobic environment and the network of hydrogen bonds provide a stable scaffold to anchor the functional and catalytic motifs. Examination of the sequence

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conservation for IMPDH representatives from the three kingdoms suggests a limited sequence conservation of the  $\alpha/\beta$  barrel core structure relative to the high level of conservation observed for residues forming the catalytic site pocket. The sequence conservation of  $\alpha/\beta$  barrel core is restricted to residues adjacent to the active site pocket and to a region representing the junction between the catalytic and CBS dimer domains.

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Several large structural and catalytic protrusions connect the  $\beta$ -strands and  $\alpha$ helices of the  $\alpha/\beta$  barrel surface. The distal face of the  $\alpha/\beta$  barrel (furthest from the IMP binding pocket) provides for entry of the N-terminus (strand \beta1). The CBS dimer domain is attached through helix  $\alpha 2$  and strand  $\beta 3$ . Connections between the remaining  $\alpha/\beta$  motifs are short (2-5 amino acid residues) and characterized by a preponderance of proline, glycine and hydrophobic residues. The C-terminal region exits from helix a8 and is located on the opposite face of the tetramer from the Nterminus.

The protrusions on the proximal face of the  $\alpha/\beta$  barrel scaffold range in size from 3-67 residues and define the character of the active site. Three of the barrel connections ( $\beta 1/\alpha 1$ ,  $\beta 6/\alpha 6$ , and  $\beta 7/\alpha 7$ ) show greater than a 50% amino acid sequence conservation for IMPDH proteins representing the three kingdoms. The  $\beta 8/\alpha 8$ protrusion is the largest (67 residues) of the proximal face motifs and contains the "finger" structure ( $\beta$ J and  $\beta$ K, FIGS. 1b, 2a), short helices  $\alpha$ I and  $\alpha$ J, strand  $\beta$ M, and regions that have a role in catalysis and that interact with other IMPDH monomers in the tetramer. This protrusion sequence is also highly conserved with regional sequence conservation of 60-80% in three distinct 10-amino acid residue segments. A distinct feature of this region is a "flap" (residues 396-419) on one edge of the active site that apparently projects into the solvent. This flap has been suggested to function by potentially folding over the catalytic pocket controlling access to and ordering the active site. (Whitby et al., 1997) This structure is similar to the active site flap involved in the catalytic mechanism of lactate dehydrogenase (Holbrook t al., 1975). In the S. pyogenes IMPDH 1.9 Å structure, 14 residues in this loop remain disordered in the presence of substrate in the active site and also in IMPDH crystals containing product, transition state analogue complexed with MPA (Sintchak et al., 1996; Whitby et al., 1997. This persistent disorder suggests that NAD binding may be

critical for structuring the flap; a suggestion supported by the resistance of this region to proteolysis acquired by NAD binding (Nimmesgern et al., 1996). This also suggests that MPA binding does not involve an interaction with this flap and does not entirely mimic NAD binding. These features suggest this flap may be important in mediating NAD binding specificity in the active site and may be responsible for some

of the kinetic differences of IMPDH enzymes from bacteria and eukaryotes.

### **CBS Dimer Domain**

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The CBS dimer domain contains residues 94-223 with an approximate size of 20x20x30 Å. A CBS domain was originally identified in cystathionine-β-synthase and proposed as a regulatory element since mutations lead to the human disease homocystinuria (Bateman, 1997). The CBS dimer domain is composed of two CBS motifs arranged approximately on a two-fold dyad axis (FIG. 5). Each CBS motif has the characteristic sheet/helix/sheet/sheet/ helix topology. This is the first reported complete structure for this domain. The CBS dimer domain does not interact with the other subunits in the active tetrameric enzyme and may not be required for activity (Sintchak *et al.*, 1996; Zhou *et al.*, 1997). Although the amino acid sequence of this domain is not as well conserved as that of the catalytic domain, all IMPDH proteins contain this domain.

In *S. pyogenes* IMPDH, these domains form a minibarrel structure that has a hydrophobic core region with hydrophilic residues on the surface. Among bacteria, the degree of amino acid conservation is highest in the E and F  $\beta$ -strands (FIG. 2a) that span the interior of the CBS dimer domain and provide a resource of hydrophobic residues. The  $\alpha$ -helices on the exterior maintain the character of this domain with hydrophilic residues on the exterior surfaces and hydrophobic residues positioned on the interior. There is a well-defined cleft between CBS motifs (approximately 15 Å in length) between the CBS motifs; this cleft may function as a potential binding site for regulatory molecules. There is not a defined role for CBS motifs in bacteria but in eukaryotic organisms they may have a role in cytoplasmic targeting, protein-protein interactions or protein regulation (Bateman, 1997). In view of these unique characteristics, it is possible that, in bacteria, this domain may possess a species-specific regulatory role.

### **Tetramer Organization**

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S. pyogenes IMPDH is a perfect tetramer with the four subunits related by a crystallographic four-fold axis. Similarly, the structures reported for Chinese hamster (Sintchak et al., 1996) and T. foetus (Whitby et al., 1997) IMPDH also display four-fold symmetry. The scope of these structures encompass the apo-enzyme and several substrate, product, and inhibitor complexes

The tetrameric structure of IMPDH is stabilized by monomeric contacts with each of the adjacent subunits. Many of these contacts originate from interactions of the N- and C-terminal regions of the adjacent monomeric units. The subunit interactions can be arranged into three groups differing in their proximity to the catalytic site and level of amino acid sequence conservation. In one group, the first 14 residues of the N-terminus project approximately 20Å from the protein core (FIG. 1a, 2b) and interact with surface residues of an adjacent IMPDH monomer. This regional contact is distal from the catalytic site and involves residues 3-12 of the N-terminus that interact with  $\beta$ -sheet residues 465-468 of an adjacent subunit. The interaction involves hydrogen bonds and salt bridges between amino acid regions that display little sequence conservation. Another loop (residues 22-30), is involved in subunit contacts with the adjacent IMPDH molecule and also forms part of the active site pocket of the adjacent subunit. This region directly contacts the αH helix that is involved in binding IMP and the  $\alpha$ 4 helix of the  $\alpha/\beta$  motif implicated in the binding of NAD (Sintchak et al., 1996). This region contains amino acid residues that are conserved in IMPDH enzymes from the three kingdoms. The sequence conservation and proximity to the active site suggests these interactions may indirectly mediate catalytic activity and account for the tetrameric character of the active enzyme. Additional subunit contacts originate from  $\beta$ -strand  $\beta$ K and residues 479-484 in an adjacent IMPDH monomer. These regions are on the exterior of the tetramer approximately 20Å from the IMP binding site and display an amino acid sequence conservation that is restricted to a specific phylogenetic group.

A feature observed in the tetramer structure is the projection of an extended region from the C-terminal face of each monomer subunit (FIG. 1b). These "fingers" are observed in all IMPDH enzymes for which structural information is available. This region of 12 amino acids forms two anti-parallel  $\beta$ -strand structures stabilized by hydrogen bonding and interactions with the  $\beta$ L region (FIG. 2a). Interestingly, in all

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IMPDH enzymes, this region contains at least two solvent-exposed hydrophobic residues. Other than a predominance of aromatic amino acids, there is little sequence conservation even within the specific phylogenetic domains. However, the  $\beta$ -strand structure of the fingers is preserved and amino acid residues at the base of the fingers are conserved for all phylogenetic groups. The conservation of this structure may have functional consequences for the interaction of the tetramer with other IMPDH complexes or cellular proteins.

### Catalytic Site and Implication for the Mechanism of Bacterial IMPDH

IMP dehydrogenase catalyzes the oxidation of inosine 5'-monophosphate to xanthosine 5'-monophosphate with the concomitant reduction of NAD to NADH. IMP is bound at one end of the barrel with the other end blocked by the  $\beta B/\beta C$  sheet (FIG. 2a). Short helices H, J, and I are structural motifs containing many of the active site residues. During the reaction the hydride is transferred from the C2 carbon of the hypoxanthine ring to NAD and an oxygen atom is substituted in the C2 position resulting in the formation of xanthosine.

The high-resolution (1.9 Å) crystal structure of S. pyogenes IMP dehydrogenase allows examination of the catalytic site in greater detail than it was possible previously. The enzyme contains the inosine monophosphate substrate bound into the pocket located near the surface of the  $\alpha/\beta$ -barrel structure. The inosine ribose and phosphate moieties are highly coordinated by protein (FIG. 3a). The sugar is in the C2'-endo-conformation and its 2'- and 3'-hydroxyls are hydrogen-bonded with the Asp343 residue as well as with a water molecule that through a water relay system connects with N3 of the hypoxanthine ring (FIG. 3a). The phosphate group is anchored in its site by a number of amino acid side chains (S308, S367 and Y390) and three main chain nitrogens (G345, G366 and S367). The remaining hydrogen-binding potential of the phosphate oxygens is realized with water molecules.

The conformation of the glycosidic torsion angle of the bound nucleotide is anti and the hypoxanthine ring interacts with the ribose and the phosphate moiety only through water mediated interactions and appears to be free to rotate around the glycosidic bond (FIG. 3a). This conformation places H2 of hypoxanthine ring (which is transferred to NAD in the reaction) in a position unobstructed by the rest of the molecule to facilitate the reaction. N1, N7 and O6 of the hypoxanthine ring are

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hydrogen bonded to the main chain carbonyl of E421 and main chain nitrogen of M393 and G394, respectively. However, N3 is not involved in an interaction with protein and only weakly with solvent. There are van der Waals contacts between the hypoxanthine ring and the Ile309 residue.

Cysteine 310 has been identified previously as a key residue in catalysis (Huete-Pérez et al., 1995; Antonino et al., 1994). The ability of the thiol residue to ionize appears to be critical for the reaction involving nucleophilic attack. The hydroxyl of T312 is in position (3.3 Å) to extract a hydrogen from C310 and therefore ionize the cysteine residue. This is consistent with mutagenesis studies that show that substitution of this residue abolishes enzyme activity (Sintchak et al., 1996). The sulfur atom is located above the plane of the hypoxanthine ring, 3.3 Å from the C2 atom, and is not covalently attached to the ring (FIG. 5c). The C310 is in a position for a nucleophilic attack on C2 carbon once the activation of the CysTEINE residue is accomplished and the orientation of the hypoxanthine ring is adjusted (it can swivel around glycosidic bond). The formation of a tetrahedral intermediate has been proposed (Xiang et al., 1997). However, the present inventions shows that IMPDH does not form a covalent bond with the substrate in the absence of the NAD cofactor. Therefore a cofactor plays not only the role of hydride acceptor but also appears to complete the structure of the catalytic pocket. Initiation of a reaction cycle requires alignment of the hypoxanthine and nicotinamide rings in near parallel fashion and positioning of the C2 of hypoxanthine ring in close contact with C4 on the beta face of nicotinamide ring (Xiang et al., 1997). This places the amide moiety of NAD near the N3 nitrogen of hypoxanthine. Such a configuration may facilitate the electron shift within the hypoxanthine ring required for formation of the thioimidate intermediate. Therefore, it appears that the cofactor may play a role in activation of the substrate. This mechanism is in striking contrast with results obtained with halogenated derivatives of IMP. Human IMPDH catalyses the dehalogenation of 2-fluoro- and 2chloroinosine 5'-monophosphate in the absence of NAD (Antonino et al., 1994). This suggests that, although the C310 activation system is in place, the reaction does not proceed with IMP because hydride is a much poorer leaving group than chlorine and fluorine and the binding of NAD is required.

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The structure of the hamster IMPDH has been reported (Sintchak et al., 1996). This structure contains the hypoxanthine ring covalently bound to C331 (equivalent to C10 in our structure) and an inhibitor MPA bound to the active site. It appears that the hamster IMPDH structure represents the covalent thioimidate intermediate of the reaction in which MPA, an uncompetitive inhibitor, prevents the hydrolysis of the thiopurine covalent intermediate as was suggested previously by Link and Straub (Link et al., 1996). Therefore MPA restricts the access of the solvent molecules and blocks subsequent steps of the reaction. This observation also suggests that the hydrolysis of the thioimidate intermediate is mediated by an activated water molecule originating from the NAD site. In the S. pyogenes IMPDH structure, two water residue were located that are potential candidates for nucleophilic attack on the thioimidate (FIG. 3b). Several residues (E421, T312, and Y450 from and adjacent subunit) in the active site pocket can act as activators of this water molecule. Because MPA can stabilize the thioimidate intermediate in the human enzyme (Sintchak et al., 1996), hydrolysis of thioimidate must be several orders of magnitude slower than the dissociation of NADH. These results are consistent with the mechanism proposed by Wang et al. (1996)

The binding of NAD to IMPDH has not been structurally characterized. However, a structure has been reported for the hamster enzyme complexed with MPA (Sintchak et al., 1996), an uncompetitive inhibitor of mammalian IMPDH enzymes. MPA has been suggested to inhibit the hamster enzyme by mimicking the nicotinamide portion of NAD and blocking access of a catalytic water molecule (Sintchak et al., 1996). Comparison of the residues involved in MPA binding in the hamster enzyme (D274, S276, N303, R322, Gl326, T333, Q441) with the equivalent residues in *S. pyogenes* IMPDH indicates these residues are largely conserved. The Asparagine, Aspartate, Glycine, and Threonine residues are identical, Threonine replaces S276 (although Serine is present in other bacterial enzymes), and Lys301 replaces the hamster R322 residue. The most significant change appears to be replacement of E421 with Q441. Interestingly, this residue is part of the active site flap that is ordered. This suggests that the NAD binding pocket is different in bacterial IMPDH, however the mechanism of IMP oxidation remains the same.

### **MATERIALS AND METHODS**

### Site-specific mutants.

To validate the role of specific residues in catalysis and to provide a basis for comparing the bacterial and mammalian enzymes, several point mutants were constructed. The sites for mutation were selected on the basis of previous studies suggesting a catalytic role for the region and supported by information derived from the *S. pyogenes* IMPDH crystal structure. One region targeted for site-specific mutagenesis was the active site flap. This flap is present in all IMPDH enzymes and is disordered in the *S. pyogenes* IMPDH structures and in the IMPDH structures from hamster and *T. foetus*. Although this region has not been previously implicated in the catalysis mechanism of IMPDH enzymes, the presence of a conserved RY(FY) motif and the similarities to the flap region in lactate dehydrogenase (Holbrook *et al.*, 1975) suggest a potential role in catalysis. Mutation of R406 to alanine in this flap region results in a complete loss of enzyme activity (Table 6) as might be expected for a residue conserved in all

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	Corresponding residue in	Reg	Relative specific	
Mutant	mammalian IMPDH	Location	Function	activity of purified
Arg406®Ala	Arg	Active site flap	Catalysis	No activity
Tyr450®Asp	Asp	Helix 8, TIM barrel	Unknown	No activity
Tyr450®Ala	Asp	Active site	Unknown	0.25
Glu421®Gln	Gln	$\alpha_J/\beta_L$ Loop	NAD binding region	1.0

IMPDH enzymes. This loss of activity confirms the importance of the active site flap in catalysis. Since there is little sequence conservation of this region, this structure is an attractive target for specific inhibitors.

The catalytic mechanism of *S. pyogenes* IMPDH involves the hydrolysis of a thioimidate intermediate that we believe is mediated by an activated water molecule originating from the NAD site. In the *S. pyogenes* IMPDH structure, we have located two water residues that are potential candidates for nucleophilic attack on the

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thioimidate. Tyrosine 450 originating from an adjacent subunit, is a residue in the active site pocket that can act as an activator of one of these water molecules. This residue is located at the noncatalytic end of a conserved helix (Helix 8) that forms the TIM barrel core. Replacement of Y450 with aspartic acid or alanine (Table 6) results in substantial loss of enzyme activity. Approximately 25% activity is retained for an alanine replacement, but substitution of aspartic acid results in a loss of enzyme activity. This region is conserved in the IMPDH enzymes, but the sequence pattern is different in bacteria and eukaryotes, suggesting this region may contribute to the differential signature of the bacterial and mammalian enzymes.

The NAD binding region (between the  $\alpha_{J/}\beta_L$  loop) was also selected as a target for site-specific mutagenesis. The selection of E421 for mutation was based on an analysis of sequence differences at residues corresponding to or near amino acids identified as MPA binding sites in human IMPDH. The conserved glutamate in bacteria is replaced with a conserved glutamine in eukaryotes. This substitution does not alter the apparent activity of *S. pyogenes* IMPDH (Table 6). This result was unexpected since replacement of the corresponding residue in the hamster enzyme (Q441) with alanine results in a significant decrease in activity (Sintchak *et al.*, 1996) Cloning and Expression of *S. Pyogenes* IMPDH

The coding region of IMPDH was amplified from *S. pyogenes* genomic DNA (provided by Dr. Michael Boyle, Medical College of Ohio, Toledo, Ohio; Genomic DNA from *S. pyogenes* is also available from the American Type Culture Collection [ATCC] as catalogue No. 700294D) using coding region-specific primers and a proofreading polymerase (Pfu). The amplified fragment was cloned into a pET23a (Novagen) expression vector and used to transform BL21(DE3)lysS bacterial cells. DNA sequencing of the expression constructs validated sequence integrity of the initiation and termination regions. Expression of Streptococcal IMPDH was induced by the addition of IPTG to a concentration of 0.5 mM.

The Streptococcal IMPDH enzyme was purified using a modification of the procedure previously described for the human enzymes (Hager et al., 1995). The modified procedure replaces the Blue Sepharose dye column with a Matrex Green resin (Millipore, Bedford, MA). Since the enzyme elutes as a broad peak from the dye column, an additional chromatographic procedure was applied to facilitated

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enzyme concentration and increase purity. Peak fractions from the dye column are diluted with 20 mM Tris-HCL, pH 7.4 and applied to a MonoQ HR10/10 FPLC column (Pharmacia, Piscataway, NJ). The column was washed with 20 mM Tris-HCl, pH 7.4, 1 mM DTT and the enzyme eluted with a linear gradient of 0.2-0.7 M NaCl in wash buffer.

Purified IMPDH from *S. pyogenes* was characterized by N-terminal sequencing and analyzed by mass spectroscopy to validate as much of the internal protein sequence as is possible. An N-terminal sequence was obtained (Yale Biotechnology Resource Center) for 19 residues corresponding to amino acids 2-20 of the predicted sequence and indicated cleavage of the N-terminal methionine as is commonly observed for proteins expressed in *E. coli*. Characterization of the purified protein also included matrix-assisted laser desorption ionization mass spectroscopy (MALDI-MS) analysis of the intact and tryptic-digested protein provided by The Biotechnology Resource Laboratory at Yale University. MALDI-MS of the intact protein indicated a molecular weight (MW) of 52,328 similar to the predicted MW of 52,657. In addition to N-terminal sequencing of the intact protein, a triptych digest of the purified protein was analyzed by MALDI-MS. This analysis provided verification of approximately 60% of the of the internal protein sequence.

Selenomethionyl IMPDH was obtained by growth of the native expression bacterium in M9 medium. Prior to induction of IMPDH expression, *de novo* methionine synthesis was suppressed by the addition of phenyalanine, valine, threonine, isoleucine, leucine, and lysine to a final concentration of 50 ug/ml. Thirty minutes later, selenomethionine was added to a final concentration of 50 ug/L and IPTG was added 0.25 mM. The induced bacteria were harvested 4-6 h after induction. The purification and crystallization of selenomethionyl IMPDH was as described for the wild-type enzyme and the presence of selenomethionine was verified by amino acid analysis of the purified protein.

### **Crystallization and Data Collection**

Crystals of IMPDH from *S. pyogenes* were grown by the hanging drop method. The reservoir solution was 0.1 M MES (pH 7.2), 1.8 M ammonium sulfate, with 1 mM IMP. The crystals grew in a few days to about 0.1 x 0.1 x 0.25 mm (maximum size). Crystals were transferred into a cryo-protectant solvent prepared by

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the addition of glycerol to the crystallization solution (final glycerol concentration (v/v), 28%). Crystals were flash cooled in liquid nitrogen for all data collections.

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Diffraction data were collected on beamline 19ID of the Structural Biology Center at the Advanced Photon Source. The approximate x-ray flux on the sample was 1 x 10 Ph/sec. Diffraction patterns from IMPDH crystals were collected at 100°K using a 3x3 mosaic CCD area detector (Westbrook et al. 1997) and data were processed by the HKL2000 (Otwinowski et al., 1997) package. Diffraction patterns of the IMPDH crystals exhibited 4/mmm symmetry. Bragg spots with indices other than (h + k + l) = 2n were systematically absent. Therefore the space group to which these crystals belong must be I422. The cell dimensions are a=b=151.49Å, c=101.67 Å,  $\alpha = \beta = \gamma = 90^{\circ}$ . Each asymmetric unit of this crystal form contains one monomer; the estimated solvent content is 55% and  $V_M = 2.79 \text{ Å}^3/\text{Da}$ .

Crystals for the MAD study were of SeMet IMPDH from S. pyogenes complexed with IMP. We recorded three data sets for a single crystal, each at a unique x-ray wavelength ( $\lambda 1 = 1.0781 \text{ Å}, \lambda_2 = 0.9793 \text{ Å}, \lambda_3 = 0.9791 \text{ Å}, \text{ Table 3}$ ). The entire time to manipulate the sample and acquire data required less than one hour. The crystal was not oriented in any special way prior to data collection. Data quality is summarized in Table 3. The high-resolution data (1.90 Å) were collected from the same crystal at wavelength  $\lambda = 1.0332$  Å. Details of the experiments and data quality are summarized in Table 3.

### **Phasing**

Phase analysis for the crystal form was initiated by carrying out molecular replacement (MR), using AMORE (Navaza et al., 1997) and the T. foetus atomic coordinates (Whitby et al., 1997) from the Protein Data Bank as a search model. The initial molecular replacement solution of this structure produced phases that were not sufficiently close to the correct values for us to interpret the structure further. However this phase set was sufficiently good to identify 6 of the 13 selenium sites in the structure. These selenium sites were refined by the method discussed by Ramakrishnan and Biou (1997), using the program MLPHARE (Otwinowski, 1991), yielding a phase set which permitted identification of two additional selenium atoms.

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Further MLPHARE refinement with 8 selenium sites produced phases that permitted location of three additional selenium sites by difference Fourier analysis. The next stage of MLPHARE refinement against 11 selenium sites produced phases with which the remaining 2 selenium sites were identified. The final round of MLPHARE phasing with all 13 selenium sites (Table 3) produced a map with which interpretation of the model was completed. Phases were improved during subsequent refinement with CNS (Holbrook *et al.*, 1975) (see below) permitting modeling of 97% of the structure.

### Model Building, and Refinement

All model building was carried out with FRODO (Jones, 1968) on an Evans and Sutherland ESV10 graphics workstation. Relative to the map obtained by molecular replacement, the MAD map obtained with six selenium sites allowed localization of IMP in the active site and corrected several errors in the catalytic domain model. The MAD map calculated with 8 selenium sites allowed modeling of the complete N-terminus (except residue 1), the C-terminus to residue 480, and the CBS dimer domain with the exception of residues 114-169. The MAD map obtained with eleven selenium sites allowed assignment of the C-terminus to residue to 490, and decreased the undefined region of the CBS dimer domain to residues 146-162. When all 13 selenium sites were used in the MAD map calculation, it was possible to model the entire molecule, with the exception of residue 1, residues 221-226, the active site flap (residues 402-415) and C-terminal residue 493. CNS refinement improved phases to allow modeling of residues 221-226 in the CBS dimer domain. This model accounts for 97% of the residues predicted from the gene sequence.

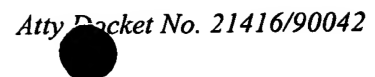
Refinement of the initial model against the MAD data was carried out using torsion-angle molecular dynamics (Rice *et al.*, 1994) and the phase restrained MLHL target (Pannu *et al.*, 1998) implemented in CNS (Holbrook *et al.*, 1975). All diffraction data (6.0-1.90 Å) were used throughout the refinement except for a 10% randomly selected test set required for cross-validation of the  $\sigma_A$  values used in the maximum likelihood target and free R calculations. A flat bulk solvent model was implemented in density modification of the initial MAD maps, with the program DM (Cowtan, 1994). At the later stages,  $\sigma_A$  phase-combined maps (Pannu *et al.*, 1998) were calculated, with model phases calculated from the MLHL refined model

combined with experimental phases. Alternate cycles of model rebuilding, positional refinement, restrained B-factor refinement, and water placement followed, decreasing the free R-factor from its initial value of 48% to 26.1% and yielding the current R-factor of 23.2% (Table 5). The model has a correlation coefficient ( $F_0$  versus  $F_c$ ) of 95% and an estimated coordinate error of 0.3Å using the SIGMAA (Read, 1986) sftware suite. Stereochemical and other refinement parameters are given in Table 4. By PROCHECK (Laskowski *et al.*, 1993) criteria, the model has 91.2% of the main chain torsion angles within the "allowed regions" of the Ramachandran plot and 8.8% within the "additional allowed regions".

## 10 Coordinates

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The coordinates of the crystalline IMPDH molecule (Table 7) have been deposited in the Brookhaven Protein Data Bank under accession number 1ZFJ.



cerevisiae

```
(SEQ 10 NO:1) -
    Table 2. Catalytic Region
                PGSIC TTRVVAGVGV Streptococcus pyogenes
  A Bacteria
              PGSIC TTRVVAGVGV
              A PGSIC TTRIVTGVGVA
5
                PGSIC TTRVVAGVGV
                                  Mycobacterium tuberculosis
              PGSIC TTRVVAGVGV
                SGSIC ITQEVLACGR Homo sand
                                  Homo sapiens
    Ecarya
                SGSIC ITQEVLACGR
10
                 SGSIC ITQEVLACGRA
                 TGSIC ITQKVMACGR
                                   prosophila melanogaster
                 SGSIC ITQEVMACGR
     Active Site Flap
15
                                                  (SEGID NO. (D)
                                                   Streptococcus pyogenes
                MA....KG SSDRYFQ.SD NAADKLVPEG ∧
     Bacteria [
                 MS.....KG SSDRYFQ.SD NAADKLVPEG
                 MK....KG SSDRYFQGSV NEANKLVPEG
                 ME.....KG SKDRYFQ... EENKKFVPEG
20
                 MRGRGGATSY SKDRYFADDA LSEDKLVPEG
                                                     tuberculosis
                                                   (SEQ IDNO .17)
               MD....KHLS SQNRYFSEAD KIK...VAQG HOMO
     Eucarya
               MD....KHLS SQNRYFSEAD KIK...VAQG
25
               MERGDAKGAA MSRYYHNEMD KMK...VAQG
               MOKTGTKGNA STSRYFSESD SVL...VAQG
               MTKG..... SDQRYLGDQT KLK...IAQG \DIC
```

M....SQGKE SGKRYLSENE AVQ...VAQG Saccharomyces

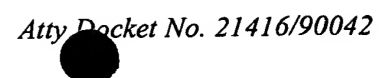


	Table 3. Crystal and MAD	Data Collection P	arameters for I	MPDH					
5	Crystal Parameters								
	Unit Cell	a = b = 151.49	$a = b = 151.49 \text{ Å, } c = 101.67 \text{ Å, } \alpha = \beta = \gamma = 90^{\circ}$						
	Space Group	I422							
	MW	53,328							
	Mol/AU	1							
10	Se-Met/AU	13							
	MAD Data Collection (Sel	Met IMPDH)							
	Oscillation Angle	1°							
	Oscillation Range	90°							
15	Exposure time/degree	5 sec							
		Edge $(\lambda_2)$	Peak $(\lambda_3)$	Remote $(\lambda_1)$					
	Wavelength (Å)	0.9793	0.9791	1.0781					
	Resolution (Å)	2.5	2.5	2.5					
20	Total observations	283910	276365	272576					
20	Unique reflections	20633	20627	20686					
	Redundancy	6.9	6.7	6.6					
	Completeness	99.7	99.7	99.6					
	R <sub>merge</sub> (%)	7.7	9.6	5.9					
25				•					
	<b>High Resolution Data Set</b>								
	Oscillation angle	1°							
	Oscillation range	90°							
	Exposure time/degree	8 sec							
30	Wavelength (Å)	1.0332							
	Resolution (Å)	30-1.90							
	Total observations	263,355							
	Unique reflections	44,921							
	Completeness (%)	96.5							
35	R <sub>merge</sub> (%)	6.8							

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### Table 4. Summary of MLPHARE Phasing

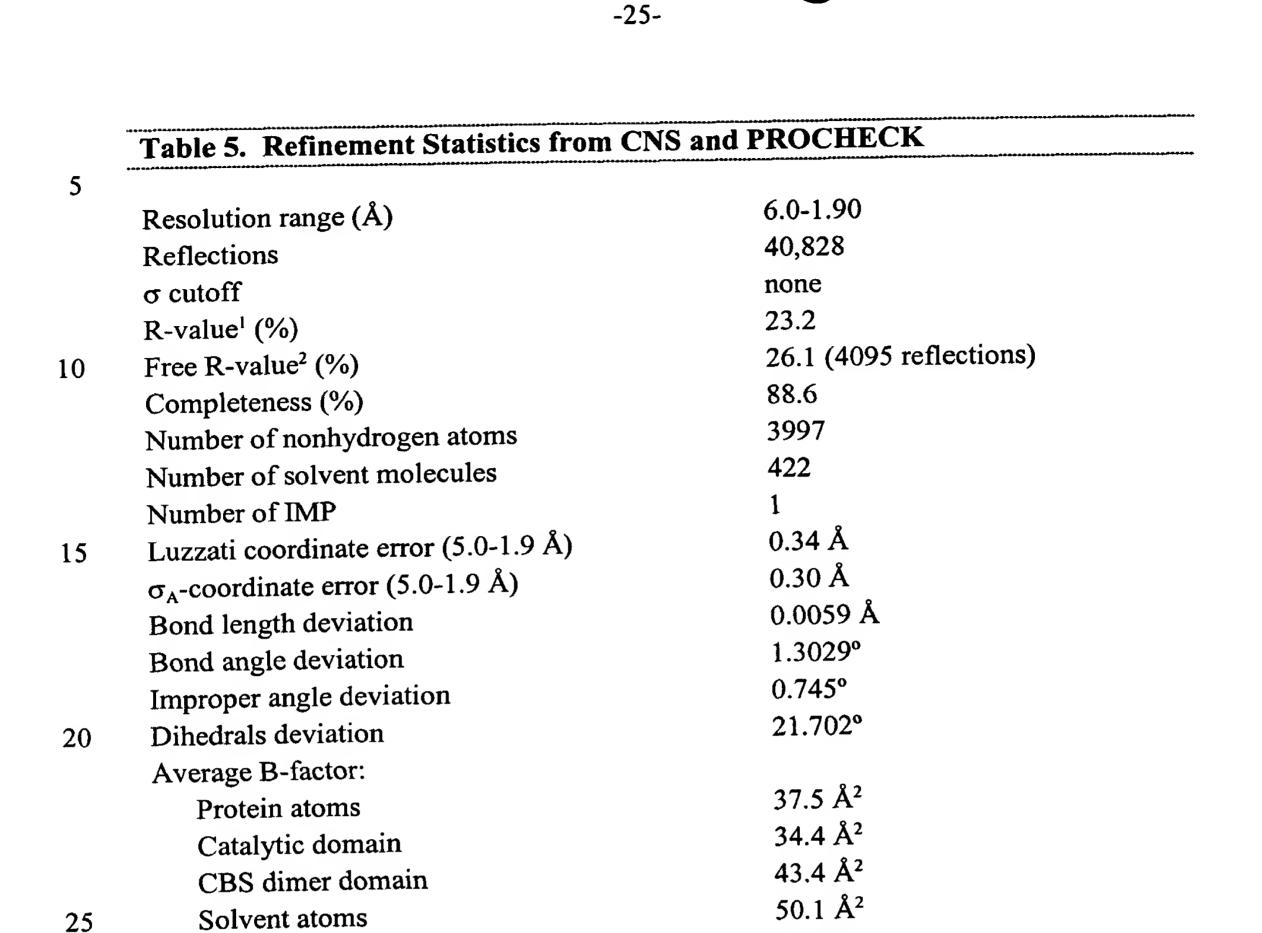
		_	Acentric		_	Centric	All		
	Resolution (Å)	No	FOM <sup>a</sup>	Phasing <sup>b</sup> power	No	FOM	Phasing power	No	FOM
10	7.27	631	0.56	1.71	231	0.51	1.71	862	0.55
	5.71	759	0.79	3.35	167	0.71	3.03	926	0.78
	4.71	1188	0.79	3.18	208	0.68	2.40	1396	0.77
	4.00	1719	0.77	2.86	242	0.65	1.92	1961	0.76
	3.48	2337	0.75	2.42	265	0.65	1.82	2602	0.74
15	3.08	3053	0.70	2.13	283	0.60	1.49	3336	0.69
	2.76	3860	0.62	1.78	270	0.54	1.09	4130	0.61
	2.5	4642	0.45	1.21	207	0.41	0.81	4849	0.45
	Total	18189	0.64	2.02	1873	0.60	1.63	2006 2	0.64

<sup>a</sup>Figure of Merit is a measure of the relative reliability of a phase based on the consistency of the MIR analysis from one derivative to the next. The maximum value is 1.0.

<sup>b</sup>MAD phasing power is defined:

 $\langle |F_{h1} - F_{hi}|^2 \rangle / \int_{b} P_{\lambda_1 \to \lambda_i}(\phi) (|F_{\lambda_1}| e^{i\phi} + F_{hi} - F_{hi}| - |F_{\lambda_i}|)^2 d\phi \rangle^{\frac{1}{2}}$  computed for individual lack-of-closure expressions between the reflections of the reference wavelength  $\lambda_1$ , its Friedel mate, and the Bijvoet pairs measured at the other wavelengths  $(F_{hi})$ .  $P_{\lambda_1 \circ \lambda_i}(\phi)$  is the corresponding phase probability distribution.

Atty Docket No. 21416/90042



91.2%

0.0%

$$^{1}$$
R-value = 
$$\frac{\left|F_{obs}\right| - \kappa \left|F_{calc}\right|}{\left|F_{obs}\right|}$$

Residues in core phi-psi regions

Residues in disallowed regions

<sup>2</sup>Free R-value is the R-value obtained for a test set of reflections (10% of the diffraction data) not used during refinement or  $\sigma_A$  calculations. 30

```
DEHYDROGENASE
HEADER
                                                    29-MAR-99
                                                                 1ZFJ
TITLE
          INOSINE MONOPHOSPHATE DEHYDROGENASE (IMPDH; EC 1.1.1.205)
TITLE
         2 FROM STREPTOCOCCUS PYOGENES
COMPND
          MOL ID: 1;
COMPND
         2 MOLECULE: INOSINE MONOPHOSPHATE DEHYDROGENASE;
COMPND
         3 CHAIN: A;
         4 FRAGMENT: CATALYTIC DOMAIN, CBS DOMAIN;
COMPND
         5 EC: 1.1.1.205;
COMPND
COMPND
         6 ENGINEERED: YES;
         7 BIOLOGICAL UNIT: TETRAMER
COMPND
SOURCE
          MOL_ID: 1;
         2 ORGANISM_SCIENTIFIC: STREPTOCOCCUS PYOGENES;
SOURCE
SOURCE
         3 EXPRESSION_SYSTEM: STREPTOCOCCUS PYOGENES;
         4 EXPRESSION_SYSTEM_STRAIN: ESCHERICHIA COLI
SOURCE
KEYWDS
          IMPDH, DEHYDROGENASE, CBS DOMAINS
EXPDTA
          X-RAY DIFFRACTION
          R. ZHANG, G. EVANS, F.J. ROTELLA, E.M. WESTBROOK, D. BENO, E. HUBERMAN,
AUTHOR
AUTHOR
         2 A.JOACHIMIAK, F.R. COLLART
JRNL
                   R. ZHANG, G. EVANS, F.J. ROTELLA, E.M. WESTBROOK, D. BENO,
            AUTH
JRNL
            AUTH 2 E.HUBERMAN, A. JOACHIMIAK, F.R. COLLART
                    CHARACTERISTICS AND CRYSTAL STRUCTURE OF BACTERIAL
            TITL
JRNL
            TITL 2 IMP DEHYDROGENASE
JRNL
JRNL
            REF
                   TO BE PUBLISHED
JRNL
            REFN
                                                                     0353
REMARK
         1
REMARK
REMARK
         2 RESOLUTION. 1.90 ANGSTROMS.
REMARK
         3 REFINEMENT.
REMARK
                          : CNS 0.3
             PROGRAM
REMARK
             AUTHORS
                          : BRUNGER, ADAMS, CLORE, DELANO, GROS, GROSSE-
REMARK
REMARK
                          : KUNSTLEVE, JIANG, KUSZEWSKI, NILGES, PANNU,
                          : READ, RICE, SIMONSON, WARREN
REMARK
REMARK
REMARK
            REFINEMENT TARGET : NULL
REMARK
REMARK
            DATA USED IN REFINEMENT.
REMARK
             RESOLUTION RANGE HIGH (ANGSTROMS) : 1.9
             RESOLUTION RANGE LOW (ANGSTROMS): 6.0
REMARK
REMARK
             DATA CUTOFF
                                     (SIGMA(F)) : 0.0
             OUTLIER CUTOFF HIGH (RMS(ABS(F))) : 986591.3
REMARK
             COMPLETENESS (WORKING+TEST) (%): 88.2
REMARK
             NUMBER OF REFLECTIONS
                                                 : 39729
REMARK
REMARK
         3
REMARK
         3 FIT TO DATA USED IN REFINEMENT.
REMARK
             CROSS-VALIDATION METHOD
REMARK
                                                : THROUGHOUT
```

#### TABLE 7

REMARK

BOND LENGTHS

```
FREE R VALUE TEST SET SELECTION
REMARK
                                               : RANDOM
REMARK
             R VALUE
                                 (WORKING SET) : 0.232
REMARK
             FREE R VALUE
                                               : 0.263
REMARK
             FREE R VALUE TEST SET SIZE
                                           (%): 10.0
             FREE R VALUE TEST SET COUNT
REMARK
                                               : 3980
             ESTIMATED ERROR OF FREE R VALUE : 0.004
REMARK
REMARK
             FIT IN THE HIGHEST RESOLUTION BIN.
REMARK
REMARK
             TOTAL NUMBER OF BINS USED
                                                  : 6
                                             (A) : 1.9
             BIN RESOLUTION RANGE HIGH
REMARK
             BIN RESOLUTION RANGE LOW
REMARK
                                              (A) : 2.01
             BIN COMPLETENESS (WORKING+TEST) (%): 70.8
REMARK
                                    (WORKING SET): 4706
REMARK
             REFLECTIONS IN BIN
             BIN R VALUE
REMARK
                                    (WORKING SET) : 0.357
             BIN FREE R VALUE
REMARK
                                                  : 0.368
             BIN FREE R VALUE TEST SET SIZE
REMARK
                                              (%):10.2
REMARK
             BIN FREE R VALUE TEST SET COUNT
                                                  : 534
             ESTIMATED ERROR OF BIN FREE R VALUE : 0.02
REMARK
REMARK
REMARK
            NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.
REMARK
             PROTEIN ATOMS
                                       : 3544
             NUCLEIC ACID ATOMS
REMARK
                                       : 0
             HETEROGEN ATOMS
REMARK
                                       : 23
REMARK
             SOLVENT ATOMS
                                       : 499
REMARK
REMARK
            B VALUES.
REMARK
             FROM WILSON PLOT
                                         (A**2) : 21.9
                               (OVERALL, A**2) : NULL
REMARK
             MEAN B VALUE
             OVERALL ANISOTROPIC B VALUE.
REMARK
              B11 (A**2) : 9.84
REMARK
REMARK
              B22 (A**2) : 9.84
              B33 (A**2) : -19.7
REMARK
REMARK
              B12 (A**2) : 0.0
REMARK
              B13 (A**2) : 0.0
              B23 (A**2) : 0.0
REMARK
REMARK
            ESTIMATED COORDINATE ERROR.
REMARK
             ESD FROM LUZZATI PLOT
REMARK
                                        (A) : 0.29
                                          (A) : 0.33
             ESD FROM SIGMAA
REMARK
                                        (A) : 5.0
             LOW RESOLUTION CUTOFF
REMARK
REMARK
            CROSS-VALIDATED ESTIMATED COORDINATE ERROR.
REMARK
                                         (A) : 0.32
REMARK
             ESD FROM C-V LUZZATI PLOT
             ESD FROM C-V SIGMAA
REMARK
                                      (A) : 0.37
REMARK
            RMS DEVIATIONS FROM IDEAL VALUES.
REMARK
                                           (A) : 0.018
```

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```
BOND ANGLES
                                     (DEGREES) : 2.2
REMARK
             DIHEDRAL ANGLES
                                     (DEGREES) : 21.8
REMARK
                                     (DEGREES) : 2.37
             IMPROPER ANGLES
REMARK
REMARK
            ISOTROPIC THERMAL MODEL : RESTRAINED
REMARK
REMARK
            ISOTROPIC THERMAL FACTOR RESTRAINTS.
REMARK
                                                     RMS
                                                            SIGMA
             MAIN-CHAIN BOND
                                           (A**2) : 1.21 ; 1.5
REMARK
                                           (A**2) : 1.92
             MAIN-CHAIN ANGLE
                                                          ; 2.0
REMARK
                                           (A**2) : 1.98
             SIDE-CHAIN BOND
REMARK
                                                          ; 2.0
                                           (A**2) : 3.02
             SIDE-CHAIN ANGLE
REMARK
                                                          ; 2.5
REMARK
REMARK
         3
            BULK SOLVENT MODELING.
REMARK
REMARK
            METHOD USED : NULL
REMARK
             KSOL
                          : NULL
             BSOL
                          : NULL
REMARK
REMARK
            NCS MODEL : NULL
REMARK
REMARK
                                                           SIGMA/WEIGHT
            NCS RESTRAINTS.
                                                     RMS
REMARK
                                              (A) : NULL
                                                          ; NULL
             GROUP
                    1
                       POSITIONAL
REMARK
                                           (A**2) : NULL
             GROUP
                       B-FACTOR
                                                          ; NULL
                    1
REMARK
REMARK
         3
            PARAMETER FILE
                                : PROTEIN REP.PARAM
REMARK
                            1
                                : WATER REP.PARAM
            PARAMETER FILE
REMARK
            PARAMETER FILE
                                : IMP.PAR
REMARK
            TOPOLOGY FILE
                                : PROTEIN. TOP
REMARK
                          1
            TOPOLOGY FILE
                                : WATER.TOP
REMARK
REMARK
            TOPOLOGY FILE
                                : IMP.TOP
         3
REMARK
         3 OTHER REFINEMENT REMARKS: BULK SOLVENT MODEL USED
REMARK
REMARK
         4 1ZFJ COMPLIES WITH FORMAT V. 2.3, 09-JULY-1998
REMARK
REMARK
         7
         7 S. PYOGENES IMPDH IS A TETRAMER WITH ITS FOUR SUBUNITS
REMARK
         7 RELATED BY A CRYSTALLOGRAPHIC FOURFOLD AXIS. EACH MONOMER
REMARK
         7 HAS A TWO-DOMAIN STRUCTURE: A CATALYTIC DOMAIN
REMARK
         7 (AMINO ACID RESIDUES 2-92 AND 224-492) FORMING THE INTERIOR
REMARK
REMARK
         7 CORE OF THE ACTIVE TETRAMERIC ENZYME AND A CBS DIMER DOMAIN
         7 (RESIDUES 93-223) PROJECTING OUTWARD FROM THE CORNERS OF
REMARK
         7 THE SQUARE. THE CBS DESIGNATION ARISES FROM THE ORIGINAL
REMARK
         7 IDENTIFICATION OF THIS FOLDING MOTIF IN THE ENZYME
REMARK
         7 CYSTATHIONINE-"BETA"-SYNTHASE [BATEMAN, A. (1997) TRENDS
REMARK
         7 BIOCHEM. SCI. 22, 12-13]. THE CBS DIMER DOMAIN, FOUND IN
REMARK
         7 IMPDH PROTEINS FROM ALL THREE KINGDOMS, IS COMPOSED OF TWO
REMARK
         7 CBS MOTIFS RELATED BY APPROXIMATE TWOFOLD SYMMETRY (RMS
REMARK
```

REMARK	7	7 DEVIATIONS BETWEEN ALPHA CARBON ATOMS: 2.7 AN	GSTROMS).										
REMARK	7	7 EACH CBS MOTIF HAS THE CHARACTERISTIC											
REMARK	7	7 SHEET/HELIX/SHEET/SHEET/HELIX TOPOLOGY. THIS	SHEET/HELIX/SHEET/SHEET/HELIX TOPOLOGY. THIS IS THE FIRST										
REMARK	7	REPORTED COMPLETE STRUCTURE OF A CBS DIMER DOMAIN, A											
REMARK	7	7 FOLDING MOTIF PROPOSED TO ACT AS A REGULATORY	ELEMENT										
REMARK	7	7 SINCE MUTATIONS LEAD TO THE HUMAN DISEASE HOM	OCYSTINURIA.										
REMARK	7	7 EACH IPMDH MONOMER CONTAINS IMP IN THE CATALY	EACH IPMDH MONOMER CONTAINS IMP IN THE CATALYTIC SITE.										
REMARK	7	7 THIS SUBSTRATE IS NOT COVALENTLY BOUND TO THE	ACTIVE SITE										
REMARK	7	CYS310 SUGGESTING THAT IMP DOES NOT FORM A COVALENT BOND											
REMARK	7	7 IN THE ABSENCE OF NAD.											
REMARK	100												
REMARK	100	THIS ENTRY HAS BEEN PROCESSED BY RCSB ON 30-M	AR-1999.										
REMARK	100	THE RCSB ID CODE IS RCSB000749.											
REMARK	200												
REMARK	200	EXPERIMENTAL DETAILS											
REMARK	200	EXPERIMENT TYPE : X-RAY DIFFR	ACTION										
REMARK	200	DATE OF DATA COLLECTION : NULL											
REMARK	200	TEMPERATURE (KELVIN): 110.0											
REMARK	200	•											
REMARK													
REMARK													
		SYNCHROTRON (Y/N) : Y											
		RADIATION SOURCE : APS											
		BEAMLINE : 19ID											
	-	X-RAY GENERATOR MODEL : NULL											
		MONOCHROMATIC OR LAUE (M/L) : M											
		WAVELENGTH OR RANGE (A): 0.9791,1.07	81										
		MONOCHROMATOR : SI(111)	02										
		OPTICS : MIRROR											
REMARK													
		DETECTOR TYPE : CCD											
		DETECTOR MANUFACTURER : ANL (SBC1)	AXA MOSAT										
		INTENSITY-INTEGRATION SOFTWARE : DENZO (HKL2)											
		DATA SCALING SOFTWARE : SCALEPACK ()	· ·										
REMARK			1114320007										
		NUMBER OF UNIQUE REFLECTIONS : 44921											
		RESOLUTION RANGE HIGH (A): 1.9											
		RESOLUTION RANGE LOW (A): 40.0											
		REJECTION CRITERIA (SIGMA(I)) : 0.0											
REMARK													
		OVERALL.											
·		COMPLETENESS FOR RANGE (%): 96.5											
		DATA REDUNDANCY : 6.2											
		R MERGE (I): 0.068											
		· · · · · · · · · · · · · · · · · · ·											
		<pre>&lt; &lt; I/SIGMA(I) &gt; FOR THE DATA SET : 6.0</pre>											
REMARK	∠00	,											

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```
REMARK 200 IN THE HIGHEST RESOLUTION SHELL.
REMARK 200 HIGHEST RESOLUTION SHELL, RANGE HIGH (A): 1.9
REMARK 200 HIGHEST RESOLUTION SHELL, RANGE LOW (A): 1.97
REMARK 200 COMPLETENESS FOR SHELL
                                      (%): 87.6
REMARK 200 DATA REDUNDANCY IN SHELL : 3.0
                               (I) : 0.319
REMARK 200 R MERGE FOR SHELL
                              (I) : NULL
REMARK 200 R SYM FOR SHELL
REMARK 200 <I/SIGMA(I) > FOR SHELL
                                           : 2.5
REMARK 200
REMARK 200 DIFFRACTION PROTOCOL: MAD
REMARK 200 METHOD USED TO DETERMINE THE STRUCTURE: MAD
REMARK 200 SOFTWARE USED: CNS, CCP4
REMARK 200 STARTING MODEL: NULL
REMARK 200
REMARK 200 REMARK: NULL
REMARK 280
REMARK 280 CRYSTAL
REMARK 280 SOLVENT CONTENT, VS (%): 49.0
REMARK 280 MATTHEWS COEFFICIENT, VM (ANGSTROMS**3/DA): NULL
REMARK 280
REMARK 280 CRYSTALLIZATION CONDITIONS: 0.1 M MES (PH 7.2), 1.8 M
REMARK 280 AMMONIUM SULFATE, 10 MM COCL2
REMARK 290
REMARK 290 CRYSTALLOGRAPHIC SYMMETRY
REMARK 290 SYMMETRY OPERATORS FOR SPACE GROUP: I 4 2 2
REMARK 290
REMARK 290
                SYMOP
                        SYMMETRY
REMARK 290
               NNNMMM
                        OPERATOR
REMARK 290
               1555
                        X,Y,Z
                 2555
                        -X,-Y,Z
REMARK 290
                 3555
                        -Y, X, Z
REMARK 290
                        Y, -X, Z
                 4555
REMARK 290
                 5555
REMARK 290
                        -X,Y,-Z
                        X, -Y, -Z
                 6555
REMARK 290
                 7555
                        Y, X, -Z
REMARK 290
                        -Y, -X, -Z
                 8555
REMARK 290
                        1/2+X, 1/2+Y, 1/2+Z
                 9555
REMARK 290
                        1/2-X, 1/2-Y, 1/2+Z
                10555
REMARK 290
                        1/2-Y, 1/2+X, 1/2+Z
                11555
REMARK 290
                        1/2+Y, 1/2-X, 1/2+Z
                12555
REMARK 290
                        1/2-X, 1/2+Y, 1/2-Z
REMARK 290
                13555
                        1/2+X, 1/2-Y, 1/2-Z
                14555
REMARK 290
                        1/2+Y, 1/2+X, 1/2-Z
                15555
REMARK 290
                        1/2-Y, 1/2-X, 1/2-Z
                16555
REMARK 290
REMARK 290
               WHERE NNN -> OPERATOR NUMBER
REMARK 290
                     MMM -> TRANSLATION VECTOR
REMARK 290
```

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REMARK	290						
REMARK		CRYSTALLO	RAI	PHIC SYMME	TRY TRANSFO	ORMATIONS	
REMARK	290	THE FOLLOW	INC	TRANSFORM	MATIONS OP	ERATE ON THE	ATOM/HETATM
REMARK	290	RECORDS IN				CRYSTALLOGR	
REMARK	290	RELATED MC	LEC	CULES.			
REMARK	290	SMTRY1	1	1.000000	0.000000	0.00000	0.00000
REMARK	290	SMTRY2	1	0.000000	1.000000	0.00000	0.00000
REMARK	290	SMTRY3	1	0.000000	0.00000	1.000000	0.00000
REMARK	290	SMTRY1	2	-1.000000	0.000000	0.00000	0.00000
REMARK	290	SMTRY2	2	0.000000	-1.000000	0.00000	0.00000
REMARK	290	SMTRY3	2	0.000000	0.000000	1.000000	0.00000
REMARK	290	SMTRY1	3	0.000000	-1.000000	0.00000	0.00000
REMARK	290	SMTRY2	3	1.000000	0.00000	0.00000	0.00000
REMARK	290	SMTRY3	3	0.000000	0.000000	1.000000	0.00000
REMARK	290	SMTRY1	4	0.000000	1.000000	0.00000	0.00000
REMARK		SMTRY2	4	-1.000000	0.00000	0.00000	0.00000
REMARK		SMTRY3	4	0.00000	0.000000	1.000000	0.00000
REMARK	_	SMTRY1	5	-1.000000	0.000000	0.000000	0.00000
REMARK		SMTRY2	5	0.000000	1.000000	0.000000	0.00000
REMARK		SMTRY3	5	0.000000	0.000000	-1.000000	0.00000
REMARK		SMTRY1	6	1.000000	0.000000	0.000000	0.00000
REMARK		SMTRY2	6	0.000000	-1.000000	0.000000	0.00000
REMARK		SMTRY3	6	0.000000	0.000000	-1.000000	0.00000
REMARK		SMTRY1	7 7	0.000000	1.000000	0.000000 0.000000	0.00000
REMARK		SMTRY2 SMTRY3	7	0.000000	0.000000	-1.000000	0.00000
REMARK REMARK		SMTRY1	8	0.000000	-1.000000	0.000000	0.00000
REMARK		SMTRY2	8	-1.000000	0.000000	0.000000	0.00000
REMARK		SMTRY3	8	0.000000	0.000000		0.00000
REMARK		SMTRY1	9	1.000000	0.000000	0.000000	75.74000
REMARK		SMTRY2	9	0.000000	1.000000	0.000000	75.74000
REMARK		SMTRY3	9	0.000000	0.000000	1.000000	50.84000
REMARK		SMTRY1	10	-1.000000	0.000000	0.000000	75.74000
REMARK	290	SMTRY2	10	0.00000	-1.000000	0.00000	75.74000
REMARK	290	SMTRY3	10	0.00000	0.000000	1.000000	50.84000
REMARK	290	SMTRY1	11	0.00000	-1.000000	0.00000	75.74000
REMARK	290	SMTRY2	11	1.000000	0.00000	0.00000	75.74000
REMARK	290	SMTRY3	11	0.000000	0.000000	1.000000	50.84000
REMARK	290	SMTRY1	12	0.000000	1.000000	0.00000	75.74000
REMARK	290	SMTRY2	12	-1.000000	0.000000	0.00000	75.74000
REMARK	290	SMTRY3	12	0.000000	0.000000	1.000000	50.84000
REMARK		SMTRY1	13	-1.000000	0.000000	0.00000	75.74000
REMARK	290	SMTRY2	13	0.000000	1.000000	0.00000	75.74000
REMARK		SMTRY3	13	0.00000	0.00000		50.84000
REMARK		SMTRY1	14	1.000000	0.00000	0.00000	75.74000
REMARK		SMTRY2	14	0.00000	-1.000000	0.00000	75.74000
REMARK	290	SMTRY3	14	0.000000	0.000000	-1.000000	50.84000

TABLE	27									
REMARK	290	SMTRY1	15 0.	000000	1.000	0000	0.0000	00	75.74	000
REMARK	290	SMTRY2		000000	0.000		0.0000		75.74	
REMARK		SMTRY3		000000		0000 -1			50.84	
REMARK		SMTRY1		000000	-1.000		0.0000		75.74	
REMARK		SMTRY2		000000			0.0000		75.74	
REMARK		SMTRY3		000000		0000 -1			50.84	
REMARK		DHIRIS	10 0.	000000	0.000	,000 -1		, ,	30.04	000
		REMARK: N	TIT.T.							
REMARK		KEPIAKK. W	ODD							
		MISSING A	TOM:							
		THE FOLLO		פשוותפ	<u> </u>	ATCCTNO	ያ <b>ለጥ</b> ር MG	z (M_MOD	יבו אווואס	ED.
		RES=RESID								_
		I=INSERTI			TIN THE	ZM T T L T C	ik; bbi	og≃seQ0	ENCE NO	MDEK;
REMARK		M RES C		ATOMS						
REMARK			A 109	CG	CD	CE	NZ			
REMARK			A 114	CG			OE2			
REMARK			A 114 A 121	CG	CD CD	OE1 NE	CZ	XTLI 1	MUO	
REMARK			A 143	CG	CD			NH1	NH2	
			A 400			NE CE	CZ	NH1	NH2	
REMARK REMARK			A 400 A 401	CG	CD CD	CE CE	NZ NZ			
REMARK	-, -		A 416	CG	OD1	ND2	1/1/21			
REMARK			A 415 A 417			CE	NZ			
REMARK			A 417 A 418	CG	CD CD1		NΔ			
			A 410 A 492	CG	CD1 C	CD2 O	CD	001	aaa	
REMARK REMARK		VAL	A 432	CA	C	O	CB	CG1	CG2	
		GEOMETRY	AND CTE	DECCHEM	TOTEL					
		SUBTOPIC:								
REMARK		SOBIOPIC:	CLUSE	CONTACT	5					
		THE FOLLO	ሴ/ ተእነረግ አጣ	OMC TITA	т <i>урс</i> г	ספר אתים	אם פע	יםעפייאד	T OOD A DUI:	T.C.
•	-	SYMMETRY		-						
		ANGSTROMS								
		SPECIAL P								
		INSTEAD O			•	•				75
	•	LOCATION								MC
REMARK		LOCATION	INDICAL	OKS AKE	NOT 1	NCHODE	יייייייייייייייייייייייייייייייייייייי	IIE CAL	COHATIO	N.S.
<b></b>		DISTANCE	CIITOFF.							
	• • •	2.2 ANGST			ርጥር እነር	ነጥ ፐ <i>እፕፕፖ</i> ር	TATMO	HVDDOG	ድእ <b>፤</b> አጥር M	c c
		1.6 ANGST								3
REMARK	_	1.0 ANGSI	ROMB PC	K CONTA	CIS II	и оп оти	ig iiibr	NGEN A	LOMB	
REMARK		ATM1 RE	e	EOI A	יישיט ב	מדפ כ	CCEOT	CCVMO	P DIS'	TANCE
				~						.10
REMARK		О НО	n ö	66	O F	ЮН	866	939	5 2	. 10
REMARK	_	CEOMEMPY	מחס מווע	אמננט ביים מיי	ፓርምስህ					
		GEOMETRY				ያለጠው አሳ	. VMME:TO	יותו הדל	т	
	_	SUBTOPIC:	CTOSE	CONTACT	э ти с	API APIA	O I MIME I I	CIC ONI	1	
REMARK	500									
PHINDPK	$\Box \land \land$	THE PAIR	ייע היאדויי	ים מו ארים	TN OT	OCE 00	ገለያጥን ረነጥ			
REMARK	_	THE FOLLO	WING AT	OMS ARE	IN CI	LOSE CO	NTACT.			

```
ATM1
                   RES C
                           SSEQI
                                   ATM2
                                          RES C
                                                 SSEOI
REMARK 500
                   HOH
                            676
                                     0
                                          HOH
                                                   508
                                                                     1.73
REMARK 500
              0
                                     OD1
                   HOH
                            641
                                          ASN A
                                                                     2.10
REMARK 500
                                                   275
              0
REMARK 500
REMARK 500 GEOMETRY AND STEREOCHEMISTRY
REMARK 500 SUBTOPIC: COVALENT BOND ANGLES
REMARK 500
REMARK 500 THE STEREOCHEMICAL PARAMETERS OF THE FOLLOWING RESIDUES
REMARK 500 HAVE VALUES WHICH DEVIATE FROM EXPECTED VALUES BY MORE
REMARK 500 THAN 4*RMSD (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN
REMARK 500 IDENTIFIER; SSEQ=SEQUENCE NUMBER; I=INSERTION CODE).
REMARK 500
REMARK 500 STANDARD TABLE:
REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1, 3 (1X, A4, 2X), 12X, F5.1)
REMARK 500
REMARK 500 EXPECTED VALUES: ENGH AND HUBER, 1991
REMARK 500
REMARK 500
            M RES CSSEQI ATM1
                                  ATM2
                                          ATM3
                                   CA
                                           C
                                               ANGL. DEV. = -6.3 DEGREES
REMARK 500
               PHE A
                        8
                            N
                                        -
REMARK 500
               THR A
                      14
                            N
                                    CA
                                           C
                                               ANGL. DEV. = -7.3 DEGREES
                                               ANGL. DEV. = 5.5 DEGREES
                                    CA
                                       _
                                           C
               ASP A
                      16
                            N
REMARK 500
                                    CA
                                           C
                                               ANGL. DEV. = 6.1 DEGREES
               VAL A
                      18
                            N
                                        _
REMARK 500
                                               ANGL. DEV. = -6.5 DEGREES
                                    CG
                                           CD
REMARK 500
               PRO A
                       22
                            CB
                                        _
                                               ANGL. DEV. = -7.0 DEGREES
                            CB
                                    CG
                                        _
                                           CD
               PRO A
                      29
REMARK 500
                                           C
                                               ANGL. DEV. = 6.8 DEGREES
               LEU A
                      42
                            N
                                    CA
                                        _
REMARK 500
                                               ANGL. DEV. = -6.8 DEGREES
                                    CA
                                           C
               ILE A
REMARK 500
                       46
                            N
                                        -
                            CB
                                    CG
                                           CD
                                               ANGL. DEV. = -6.7 DEGREES
               PRO A
                       47
REMARK 500
                                           C
                                               ANGL. DEV. = 10.1 DEGREES
                                    CA
REMARK 500
               THR A
                       57
                            N
                                               ANGL. DEV. =-10.6 DEGREES
               GLY A
                                    \mathsf{C}\mathsf{A}
                                        -
                                           C
                       58
                            \mathbf{N}
REMARK 500
                                    CA
                                           C
                                               ANGL. DEV. = -5.8 DEGREES
               GLU A
                       86
                            N
                                        -
REMARK 500
                                    CA
                                           C
                                               ANGL. DEV. = -6.1 DEGREES
               ASN A
                       95
                            N
REMARK 500
                                               ANGL. DEV. = -6.4 DEGREES
                                    CA
                                           C
               ILE A 99
                            N
REMARK 500
                                               ANGL. DEV. = -6.8 DEGREES
               PRO A 101
                                    CG
                                           CD
                            CB
REMARK 500
                                               ANGL. DEV. = -7.3 DEGREES
               PHE A 102
                                    CA
                                        _
                                           C
                            N
REMARK 500
                                               ANGL. DEV. = -7.2 DEGREES
                                    CG
                                           CD
               PRO A 106
                            CB
                                        _
REMARK 500
                                               ANGL. DEV. = 6.4 DEGREES
                                    CB
                                           CG
               LEU A 116
                            CA
REMARK 500
                                               ANGL. DEV. = -6.9 DEGREES
                                           CD
                            CB
                                    CG
                                        -
               PRO A 126
                                 -
REMARK 500
                                               ANGL. DEV. = -6.0 DEGREES
                                           C
                                    CA
                                        -
REMARK 500
               LYS A 135
                            N
               PRO A 154
                                               ANGL. DEV. = -7.5 DEGREES
                                    CG
                                           CD
                            CB
                                        -
REMARK 500
                                               ANGL. DEV. = -7.0 DEGREES
               HIS A 158
                                    CA
                                           C
                            N
REMARK 500
                                               ANGL. DEV. = -5.6 DEGREES
                                    CA
               GLU A 162
                                           C
                            N
                                        _
REMARK 500
                                    CA
                                                ANGL. DEV. = -5.8 DEGREES
                                           C
               HIS A 163
                            N
REMARK 500
                                                ANGL. DEV. = -6.1 DEGREES
                                           C
                            N
                                    CA
REMARK 500
               THR A 171
                                                ANGL. DEV. = -7.2 DEGREES
                                    CG
                                           CD
                            CB
                                        -
REMARK 500
               PRO A 189
                                                ANGL. DEV. = 6.3 DEGREES
                                    CA
                                           C
                                        _
               PRO A 213
                            И
REMARK 500
                                                ANGL. DEV. = -7.9 DEGREES
                                           CD
                                    CG
               PRO A 213
                            CB
REMARK 500
                                    CA
                                           C
                                                ANGL. DEV. = -6.8 DEGREES
                            N
               ALA A 216
REMARK 500
```

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### TABLE 7

REMARK 525 SOLVENT

```
C
                                                ANGL. DEV. = -7.0 DEGREES
                                    CA
REMARK 500
               VAL A 229
                            Ν
                                 -
               ILE A 252
                                           C
REMARK 500
                                    CA
                                                ANGL. DEV. = -8.2 DEGREES
                            N
               PRO A 274
                                           C
                                                              6.7 DEGREES
REMARK 500
                            N
                                    CA
                                                ANGL. DEV. =
               PRO A 274
                                                ANGL. DEV. = -7.9 DEGREES
                                    CG
                                           CD
REMARK 500
                            CB
                                        _
               ILE A 279
                                    CA
                                           C
                                                ANGL. DEV. = -9.5 DEGREES
                            N
                                        _
REMARK 500
                                           C
               ALA A 280
                                                ANGL. DEV. =
                                                               7.8 DEGREES
REMARK 500
                            N
                                    CA
               GLY A 281
                                           C
                                                ANGL. DEV. =
                                    CA
                                                               6.4 DEGREES
REMARK 500
                            N
               GLY A 281
                                                               6.2 DEGREES
                            C
                                    N
                                        _
                                           CA
                                                ANGL. DEV. =
REMARK 500
                                                ANGL. DEV. =
                                                               6.9 DEGREES
                                           C
               ASN A 282
                                    CA
REMARK 500
                            N
                                        _
               ILE A 283
                                           C
                                                ANGL. DEV. = -6.3 DEGREES
                                    CA
REMARK 500
                            Ν
                                           C
                                                ANGL. DEV. =
                                                               9.1 DEGREES
REMARK 500
               GLY A 305
                            N
                                    CA
                                           C
               PRO A 306
                                                ANGL. DEV. =
                                                               6.5 DEGREES
                                    CA
REMARK 500
                            N
               PRO A 306
                                    CG
                                           CD
                                                ANGL. DEV. = -6.4 DEGREES
                            CB
                                        _
REMARK 500
                                                ANGL. DEV. = -7.5 DEGREES
REMARK 500
                                           CD
               PRO A 321
                            CB
                                    CG
                                        _
               GLN A 322
                                           C
                                    CA
                                                ANGL. DEV. =
                                                              9.4 DEGREES
                            N
REMARK 500
               THR A 339
                                           C
                                                ANGL. DEV. =
                                                               8.8 DEGREES
                                    CA
                                        _
REMARK 500
                            Ν
               ILE A 341
                                           C
                                                ANGL. DEV. = -5.8 DEGREES
REMARK 500
                                    CA
                            N
                                        _
                                                ANGL. DEV. = -6.2 DEGREES
                                    CA
                                           C
REMARK 500
               ALA A 342
                            N
               ASP A 343
                                           C
                                                ANGL. DEV. =
                                                               5.8 DEGREES
REMARK 500
                                    CA
                            \mathbf{N}
                                           C
                                                ANGL. DEV. =
                                                               9.5 DEGREES
REMARK 500
                                    CA
                                        _
               GLY A 344
                            Ν
               ALA A 370
                                           C
                                                ANGL. DEV. = 11.1 DEGREES
REMARK 500
                            N
                                    CA
                                        _
                                                             6.6 DEGREES
                                           C
REMARK 500
                                                ANGL. DEV. =
               GLU A 374
                                    CA
                            Ν
                                        -
               PRO A 376
                                    CG
                                           CD
                                                ANGL. DEV. = -7.0 DEGREES
                            CB
REMARK 500
                                                ANGL. DEV. = -5.8 DEGREES
                                           C
REMARK 500
               LYS A 388
                            N
                                    CA
                                        _
               TYR A 390
                                           C
REMARK 500
                                    CA
                                                ANGL. DEV. = -7.0 DEGREES
                                        -
                            N
                                 _
                                                ANGL. DEV. = -7.1 DEGREES
               PRO A 420
                            CB
                                    CG
                                           CD
REMARK 500
                                        _
                                                ANGL. DEV. =
               GLU A 461
                                           C
                                                               7.6 DEGREES
                                    CA
REMARK 500
                            Ν
                                        _
                                                               8.8 DEGREES
                                           C
                                                ANGL. DEV. =
REMARK 500
               ASN A 462
                                    CA
                                        _
                            N
                                                ANGL. DEV. = -9.7 DEGREES
                                           C
               VAL A 466
                            N
                                    CA
                                        _
REMARK 500
                                                ANGL. DEV. = -5.6 DEGREES
                                    CA
                                           C
REMARK 500
               PRO A 478
                            N
                                        -
                                                ANGL. DEV. = -7.4 DEGREES
               PRO A 478
                            CB
                                    CG
                                           CD
REMARK 500
                            CB - CG - CD ANGL. DEV. = -7.9 DEGREES
REMARK 500
               PRO A 488
REMARK 500
REMARK 500 GEOMETRY AND STEREOCHEMISTRY
REMARK 500 SUBTOPIC: TORSION ANGLES
REMARK 500
REMARK 500 TORSION ANGLES OUTSIDE THE EXPECTED RAMACHANDRAN REGIONS:
REMARK 500 (M=MODEL NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER;
REMARK 500 SSEQ=SEQUENCE NUMBER; I=INSERTION CODE).
REMARK 500 STANDARD TABLE:
REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1, 4X, F7.2, 3X, F7.2)
REMARK 500
REMARK 500 M RES CSSEQI
                                              PHI
                                   PSI
REMARK 500
                                 55.58
                                          166.71
               SER A 491
REMARK 500
REMARK 525
```

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```
REMARK 525 THE FOLLOWING SOLVENT MOLECULES LIE FARTHER THAN EXPECTED
REMARK 525 FROM THE PROTEIN OR NUCLEIC ACID MOLECULE AND MAY BE
REMARK 525 ASSOCIATED WITH A SYMMETRY RELATED MOLECULE (M=MODEL
REMARK 525 NUMBER; RES=RESIDUE NAME; C=CHAIN IDENTIFIER; SSEQ=SEQUENCE
REMARK 525 NUMBER; I=INSERTION CODE):
REMARK 525
REMARK 525
           M RES CSSEQI
                              DISTANCE = 8.02 ANGSTROMS
REMARK 525
           0 HOH
                    639
REMARK 525
                    730
                              DISTANCE =
                                          6.58 ANGSTROMS
           0 HOH
                    861
                              DISTANCE =
                                          5.23 ANGSTROMS
REMARK 525
            0 HOH
                              DISTANCE =
                                          5.97 ANGSTROMS
REMARK 525
            0 HOH
                    971
REMARK 525
           0 HOH
                    983
                              DISTANCE =
                                          6.42 ANGSTROMS
                                          6.03 ANGSTROMS
            0 HOH
                    999
                              DISTANCE =
REMARK 525
                              DISTANCE =
                                          5.20 ANGSTROMS
            0 НОН
                   1021
REMARK 525
REMARK 525
            0 HOH
                   1079
                              DISTANCE =
                                          5.11 ANGSTROMS
                                          5.09 ANGSTROMS
REMARK 525
            0 HOH
                   1095
                              DISTANCE =
                                          6.40 ANGSTROMS
REMARK 525
            0 HOH
                   1136
                              DISTANCE =
                                          6.38 ANGSTROMS
REMARK 525
            0 HOH
                   1167
                              DISTANCE =
REMARK 525
                                          5.10 ANGSTROMS
            о нон
                   1173
                              DISTANCE =
            0 НОН
                   1181
                              DISTANCE =
                                          6.76 ANGSTROMS
REMARK 525
                                          5.68 ANGSTROMS
            о нон
                   1190
                              DISTANCE =
REMARK 525
                               DISTANCE =
            о нон
                   1207
                                          8.81 ANGSTROMS
REMARK 525
                              DISTANCE =
REMARK 525
            о нон
                   1216
                                           7.22 ANGSTROMS
REMARK 525
                                          6.43 ANGSTROMS
            0 HOH
                   1227
                              DISTANCE =
REMARK 525
                                          7.10 ANGSTROMS
            0 HOH
                   1282
                              DISTANCE =
REMARK 525
                              DISTANCE =
                                          6.33 ANGSTROMS
            0 HOH
                   1302
                                          5.01 ANGSTROMS
REMARK 525
            0 HOH
                   1309
                              DISTANCE =
REMARK 525
            0 HOH
                   1314
                              DISTANCE =
                                          5.92 ANGSTROMS
            0 НОН
                                          5.40 ANGSTROMS
                   1316
                               DISTANCE =
REMARK 525
                               DISTANCE =
                                           5.08 ANGSTROMS
REMARK 525
            о нон
                   1328
REMARK 525
                              DISTANCE =
                                          6.15 ANGSTROMS
            0 НОН
                   1350
REMARK 525 0 HOH 1351
                              DISTANCE = 6.96 ANGSTROMS
REMARK 525 0 HOH 1352
                             DISTANCE = 7.28 ANGSTROMS
REMARK 800
REMARK 800 SITE
REMARK 800 SITE IDENTIFIER: ASC
REMARK 800 SITE DESCRIPTION:
REMARK 800 ACTIVE SITE CYSTEINE
REMARK 800
                                P50099 IMDH_STRPY 2
DBREF 1ZFJ A 2 491 SWS
                                                               491
SEQADV 1ZFJ TYR A 387 SWS 1ZFJ A
                                       PHE
                                            387 CONFLICT
                        SWS
                             1ZFJ A
                                       GLY 402 GAP IN THE PDB ENTRY
SEQADV 1ZFJ
                                      SER
                                            403 GAP IN THE PDB ENTRY
                             1ZFJ A
SEOADV 1ZFJ
                        SWS
                                       SER 404 GAP IN THE PDB ENTRY
                        SWS
                             1ZFJ_A
SEQADV 1ZFJ
                                      ASN 405 GAP IN THE PDB ENTRY
                        SWS 1ZFJ A
SEQADV 1ZFJ
                                            406 GAP IN THE PDB ENTRY
                        SWS
                             1ZFJ A
                                       ARG
SEQADV 1ZFJ
SEQADV 1ZFJ
                                            407 GAP IN THE PDB ENTRY
                                       TYR
                        SWS
                             1ZFJ_A
```

SEQADV	1ZFJ			SWS	3 1	ZFJ_	A	PHE	4 (	08 G	AP I	N TH	E PDE	EN'	ΓRΥ
SEQADV	1ZFJ			SWS	5 1	ZFJ_	A	GLN	4 (	)9 G	AP I	N TH	E PDE	EN'	rry
SEQADV	1ZFJ			SWS	5 1	ZFJ_	A	GLY	4:	10 G	AP I	N TH	E PDE	EN'	rry
SEQADV	1ZFJ			SWS	5 1	ZFJ_	A	SER	4:	L1 G	AP I	N TH	E PDE	EN'	rry
SEQADV	1ZFJ			SWS	5 1	ZFJ_	Ą	VAL	4:	12 G	AP I	N TH	E PDE	EN'	ľRY
SEQADV	1ZFJ			SWS	3 1	ZFJ_	A	ASN	4:	L3 G	AP II	N TH	E PDE	EN'	rry
SEQADV	1ZFJ			SWS		ZFJ_		GLU	4:	L4 G	AP II	N TH	E PDE	EN'	rry
SEQADV				SWS		ZFJ_		ALA	4:	L5 G	AP I	N TH	E PDE	ENT	rry
SEQADV						ZFJ_		MET			NGIN]				
SEQADV				SWS		ZFJ_		MET			NGIN]				
SEQADV				SWS		ZFJ_Z		MET			NGIN				
SEQADV				SWS		ZFJ_Z		MET			NGIN				
SEQADV				SWS		ZFJ_/		MET			NGIN				
SEQADV				SWS		ZFJ_A		MET			NGIN				
SEQADV						ZFJ_1		MET			NGIN				
SEQADV				SWS		ZFJ_		MET			NGIN				
SEQADV				SWS		ZFJ_/		MET			NGIN				
SEQADV				SWS		ZFJ_A		MET			NGIN				
SEQADV				SWS		ZFJ_A		MET			NGIN				
SEQADV				SWS		ZFJ_A		MET			NGIN				
SEQADV				SWS		ZFJ_A		MET			NGIN		GLY	mvp.	mun.
SEQRES	1 A			_				_							
SEQRES	2 A			ASP PRO	ASP				LEU		ALA	LYS		ALA	VAL
SEQRES	3 A			LEU		LEU			PRO	ILE			ALA		
SEQRES	4 A 5 A				VAL								ALA		
SEQRES SEQRES	5 A												ASN		SER
SEQRES	7 A												VAL		ARG
SEQRES	8 A										PRO			LEU	THR
SEQRES	9 A										GLU			MSE	GLN
SEQRES	10 A				ARG				VAL			VAL		THR	
SEQRES	11 A			ASN									ASN		ASP
SEQRES	12 A			ARG							ALA			SER	
SEQRES	13 A			MSE									ALA		GLY
SEQRES	14 A			ASP									HIS		HIS
SEQRES	15 A		ARG	ILE							ASP			GLY	
SEQRES	16 A	477	LEU	SER	GLY	LEU	ILE	THR	ILE	LYS	ASP	ILE	GLU	LYS	VAL
SEQRES	17 A	477	ILE	GLU	PHE	PRO	HIS	ALA	ALA	LYS	ASP	GLU	PHE	GLY	ARG
SEQRES	18 A	477	LEU	LEU	VAL	ALA	ALA	ALA	VAL	GLY	VAL	THR	SER	ASP	THR
SEQRES	19 A	477	PHE	GLU	ARG	ALA	GLU	ALA	LEU	PHE	GLU	ALA	GLY	ALA	ASP
SEQRES	20 A	477	ALA	ILE	VAL	ILE	ASP	THR	ALA	HIS	GLY	HIS	SER	ALA	GLY
SEQRES	21 A	477	VAL	LEU	ARG	LYS	ILE	ALA	GLU	ILE	ARG	ALA	HIS	PHE	PRO
SEQRES	22 A	477	ASN	ARG	THR	LEU	ILE	ALA	GLY	ASN	ILE	ALA	THR	ALA	GLU
SEQRES	23 A	477	GLY	ALA	ARG	ALA	LEU	TYR	ASP	ALA	GLY	VAL	ASP	VAL	VAL
SEQRES	24 A	477	LYS	VAL	GLY	ILE	GLY	PRO	GLY	SER	ILE	CYS	THR	THR	ARG
SEQRES	25 A	477	VAL	VAL	ALA	GLY	VAL	GLY	VAL	PRO	GLN	VAL	THR	ALA	ILE
SEQRES	26 A	477	TYR	ASP	ALA	ALA	ALA	VAL	ALA	ARG	GLU	TYR	GLY	LYS	THR

4

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#### TABLE 7

```
SEQRES
           27 A
                  477
                       ILE ILE ALA ASP GLY GLY ILE LYS TYR SER GLY ASP ILE
                       VAL LYS ALA LEU ALA ALA GLY GLY ASN ALA VAL MSE LEU
  SEQRES
           28 A
                  477
                       GLY SER MSE PHE ALA GLY THR ASP GLU ALA PRO GLY GLU
  SEQRES
           29 A
                  477
  SEQRES
                       THR GLU ILE TYR GLN GLY ARG LYS TYR LYS THR TYR ARG
           30 A
                  477
  SEQRES
                       GLY MSE GLY SER ILE ALA ALA MSE LYS LYS ASN LYS LEU
           31 A
                  477
  SEQRES
           32 A
                       VAL PRO GLU GLY ILE GLU GLY ARG VAL ALA TYR LYS GLY
                  477
                       ALA ALA SER ASP ILE VAL PHE GLN MSE LEU GLY GLY ILE
  SEQRES
           33 A
                  477
  SEQRES
                       ARG SER GLY MSE GLY TYR VAL GLY ALA GLY ASP ILE GLN
                  477
           34 A
  SEQRES
           35 A
                  477
                       GLU LEU HIS GLU ASN ALA GLN PHE VAL GLU MSE SER GLY
  SEQRES
           36 A
                 477
                       ALA GLY LEU ILE GLU SER HIS PRO HIS ASP VAL GLN
                       THR ASN GLU ALA PRO ASN TYR SER VAL (SEQ 10 NO. 23)
SEQRES
           37 A
                  477
                            MET
  MODRES 1ZFJ MSE A
                        53
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                            MET
                        61
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                            MET
                                  SELENOMETHIONINE
                        78
  MODRES 1ZFJ MSE A
                                  SELENOMETHIONINE
                            MET
                       117
  MODRES 1ZFJ MSE A
                       145
                            MET
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                       159
                            MET
                                  SELENOMETHIONINE
  MODRES 12FJ MSE A
                       364
                            MET
                                  SELENOMETHIONINE
  MODRES 12FJ MSE A
                       368
                            MET
                                  SELENOMETHIONINE
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                       393
                            MET
  MODRES 1ZFJ MSE A
                       399
                            MET
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                       440
                            MET
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                       448
                            MET
                                  SELENOMETHIONINE
  MODRES 1ZFJ MSE A
                            MET
                       468
                                  SELENOMETHIONINE
  HET
          MSE
               A
                   53
                            8
          MSE
                   61
                            8
  HET
               Α
          MSE
               Α
                   78
                            8
  HET
               A 117
                            8
  HET
          MSE
          MSE
               A 145
                            8
  HET
  HET
          MSE
               A 159
  HET
          MSE
               A 364
                            8
  HET
               A 368
                            8
          MSE
               A 393
  HET
          MSE
               A 399
  HET
          MSE
                            8
  HET
          MSE
               A 440
  HET
          MSE
               A 448
                            8
  HET
          MSE
               A 468
                  500
                           23
  HET
          IMP
              MSE SELENOMETHIONINE
  HETNAM
              IMP INOSINE-5'-MONOPHOSPHATE
  HETNAM
  FORMUL
                       13 (C5 H11 N1 O2 SE1)
               MSE
                       C10 H13 N4 O8 P1
  FORMUL
               IMP
                      *499(H2 O1)
  FORMUL
               HOH
  HELIX
                1 ASN A
                            3
                                THR A
  HELIX
                2 PHE A
                               ASP A
                                        17
                                            5
                           15
  HELIX
                 3 PRO A
                           29
                               GLU A
                                        31
```

4 SER A

HELIX

ARG A

59

5

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HELIX	5 5	ILE A 80	SER A	93 1					14
HELIX	6 6	VAL A 110	ARG A	119 1					10
HELIX	7 7	ASN A 142	PHE A	147 5					6
HELIX	8 8	LEU A 173	GLU A	182 1					10
HELIX	9 9	ILE A 203	GLU A	211 1					9
HELIX	10 10	THR A 235	ALA A	245 1					11
HELIX	11 11	ALA A 260	HIS A	272 1					13
HELIX	12 12	ALA A 286	ASP A	294 1					9
HELIX	13 13	THR A 312	VAL A	315 1					4
HELIX	14 14	GLN A 322	GLU A	335 1					14
HELIX	15 15	SER A 349	ALA A	357 1					9
HELIX	16 16	ALA A 433	VAL A	451 1					19
HELIX	17 17	ILE A 456	ASN A	462 1					7
HELIX	18 18	GLY A 470	SER A	476 1					7
SHEET	1 A	3 PHE A 10	3 THR A	105 0					
SHEET	2 A	3 GLY A 12	4 VAL A	128 1	N PRO	A 126	O LEU	A 104	
SHEET	3 A	3 LEU A 13	6 THR A	141 -1	N ILE	A 140	O VAL	A 125	
SHEET	1 B	3 THR A 16	6 ALA A	168 0					
SHEET	2 B				N PRO	A 189	O ALA	A 167	
SHEET	3 B				N ILE	A 201	O LEU	A 188	
SHEET	1 C			A 229 0					
SHEET	2 C				N ALA	A 249	O ALA	A 227	
SHEET	3 C					A 279	O ILE	A 250	
SHEET	4 C					A 299	O ALA	A 280	
SHEET	5 C					A 339		A 300	
SHEET	1 D								
SHEET	2 D					A 387	O GLU	086 A	
SHEET	3 D					A 427		A 388	
SHEET	1 E		6 ALA A						
SHEET	2 E		2 LEU A		N LEU	A 44	O THR	A 36	
SITE	1 ASC								
CRYST1	151.48		101.680	90.00	90.00	90.00	[422	16	
ORIGX1				0.00000		0.00000			
ORIGX2				0.00000		0.00000			
ORIGX3				.000000		0.00000			
SCALE1				0.00000		0.00000			
SCALE2				0.00000		0.00000			
SCALE3				0.009835		0.00000			
ATOM	1 N				51.770	46.774	1.00 44	.22	N
ATOM		A SER A			51.755	45.463	1.00 42		C
ATOM	3 C				53.170	44.968	1.00 42		C
ATOM	4 0				53.941	45.653	1.00 42		o
ATOM		B SER A		30.340	50.980	45.597	1.00 42		Ċ
		G SER A		79.625	51.016	44.380	1.00 45		ō
MOTA	7 N			31.847	53.515	43.782	1.00 40		N
ATOM		ASN A		31.598	54.837	43.702	1.00 37		C
ATOM				30.203	54.939	42.620	1.00 37		G
ATOM	9 C	: ASN A	3 8		JZ . JJJ	74.040	1.00 57	• • • •	•

# TABLE 7 ATOM

ATOM	10	0	ASN	A	3	79.891	55.902	41.928	1.00 37.67	0
ATOM	11	CB	ASN	A	3	82.617	55.192	42.146	1.00 36.54	C
ATOM	12	CG	ASN	A	3	84.019	55.274	42.674	1.00 35.28	С
ATOM	13	OD1	ASN	A	3	84.244	55.741	43.787	1.00 35.10	0
ATOM	14	ND2	ASN	A	3	84.984	54.853	41.861	1.00 33.91	N
ATOM	15	N	TRP	Α	4	79.364	53.940	42.857	1.00 36.95	N
ATOM	16	CA	TRP	Α	4	78.018	53.998	42.322	1.00 36.01	С
MOTA	17	C	TRP	A	4	77.300	55.217	42.888	1.00 36.93	C
ATOM	18	0	TRP	A	4	76.428	55.782	42.236	1.00 37.98	0
ATOM	19	CB	TRP	Α	4	77.240	52.718	42.643	1.00 33.82	C
ATOM	20	CG	TRP	A	4	75.814	52.835	42.261	1.00 30.69	C
ATOM	21	CD1	TRP	A	4	74.805	53.326	43.029	1.00 29.45	С
ATOM	22	CD2	TRP	Α	4	75.257	52.614	40.956	1.00 29.97	C
ATOM	23	NE1	TRP	A	4	73.652	53.434	42.286	1.00 30.83	N
ATOM	24	CE2	TRP	A	4	73.902	53.007	41.010	1.00 29.48	C
ATOM	25	CE3	TRP	A	4	75.775	52.129	39.747	1.00 28.30	C
ATOM	26	CZ2	TRP	A	4	73.057	52.928	39.901	1.00 28.18	С
ATOM	27	CZ3	TRP	Α	4	74.936	52.052	38.646	1.00 27.32	C
ATOM	28	CH2	TRP	Α	4	73.592	52.450	38.731	1.00 28.03	C
ATOM	29	N	ASP		5	77.673	55.632	44.096	1.00 37.49	N
ATOM	30	CA	ASP		5	77.054	56.804	44.705	1.00 38.18	C
ATOM	31	С	ASP		5	77.778	58.093	44.327	1.00 37.52	С
ATOM	32	0	ASP		5	77.324	59.189	44.643	1.00 37.71	0
ATOM	33	CB	ASP		5	76.992	56.650	46.221	1.00 40.78	C
ATOM	34	CG	ASP		5	76.071	55.524	46.643	1.00 44.01	C
ATOM	35	OD1		A	5	74.886	55.540	46.232	1.00 43.98	0
ATOM	36	OD2	ASP		5	76.531	54.626	47.386	1.00 46.63	0
ATOM	37	N	THR		6	78.906	57.946	43.643	1.00 36.92	N
ATOM	38	CA	THR		6	79.696	59.084	43.169	1.00 35.85	C
ATOM	39	C	THR		6	79.911	58.884	41.662	1.00 33.68	C
ATOM	40	0	THR		6	80.992	59.143	41.136	1.00 33.60	0
ATOM	41	CB	THR		6	81.068	59.154	43.867	1.00 36.00	C
ATOM	42	OG1	THR		6	80.881	59.204	45.283	1.00 38.44	O C
ATOM	43	CG2 N	THR LYS		6 7	81.810 78.863	60.400 58.404	43.444	1.00 37.48 1.00 30.86	N
ATOM	44 45	CA	LYS		7	78.879	58.132	40.992 39.559	1.00 30.88	C
ATOM ATOM		CA	LYS		7	79.207	59.389	38.739	1.00 28.48	C
ATOM	46 47	0	LYS		7	79.990	59.341	37.791	1.00 28.89	0
ATOM	48	CB	LYS		7	77.523	57.553	39.153	1.00 25.62	C
ATOM	49	CG	LYS		7	77.415	57.120	37.712	1.00 23.52	C
ATOM	50	CD	LYS		7	78.423	56.039	37.712	1.00 23.54	C
ATOM	51	CE	LYS		7	78.212	54.773	38.186	1.00 23.34	C
ATOM	52	NZ	LYS		7	79.139	53.678	37.755	1.00 22.70	N
ATOM	53	N	PHE		8	78.603	60.514	39.102	1.00 22.25	N
ATOM	54	CA	PHE		8	78.860	61.781	38.420	1.00 20.00	C
ATOM	55	C	PHE		8	79.805	62.533	39.339	1.00 30.23	C
ATOM	56	0	PHE		8	79.392	63.206	40.278	1.00 32.04	0
	- <b>-</b>	_	• <b>- •</b>		-					-

MOTA	57	CB	PHE	A	8	77.542	62.513	38.205	1.00	29.85	С
ATOM	58	CG	PHE	Α	8	76.578	61.731	37.370	1.00	30.55	C
ATOM	59	CD1	PHE	A	8	76.775	61.602	36.002	1.00	29.92	С
ATOM	60	CD2	PHE	A	8	75.544	61.019	37.963	1.00	30.12	C
ATOM	61	CE1	PHE	A	8	75.958	60.770	35.241	1.00	30.48	C
MOTA	62	CE2	PHE	Α	8	74.730	60.193	37.209	1.00	28.97	C
ATOM	63	CZ	PHE	A	8	74.937	60.065	35.850	1.00	29.91	C
ATOM	64	N	LEU	A	9	81.090	62.394	39.049	1.00	32.70	N
MOTA	65	CA	LEU	Α	9	82.135	62.968	39.870	1.00	34.57	C
MOTA	66	С	LEU		9	82.222	64.485	39.903	1.00	36.07	C
ATOM	67	0	LEU		9	82.161	65.092	40.978	1.00	37.88	0
ATOM	68	CB	LEU	A	9	83.482	62.380	39.451	1.00	33.71	C
MOTA	69	CG	LEU	A	9	84.593	62.591	40.469	1.00	32.58	C
ATOM	70	CD1	LEU		9	84.139	62.012	41.789	1.00	32.56	C
MOTA	71	CD2	LEU		9	85.872	61.928	40.002	1.00	31.98	C
MOTA	72	N	LYS		10	82.377	65.104	38.740	1.00	35.80	N
MOTA	73	CA	LYS		10	82.482	66.554	38.696	1.00	35.27	C
ATOM	74	С	LYS	A	10	82.560	67.096	37.284	1.00	34.79	C
MOTA	75	0	LYS		10	82.440	66.341	36.321	1.00	35.02	0
ATOM	76	CB	LYS		10	83.699	67.008	39.514	1.00	35.85	С
ATOM	77	CG	LYS		10	84.974	66.179	39.333		33.88	C
ATOM	78	CD	LYS		10	85.554	66.275	37.947		33.98	С
ATOM	79	CE	LYS		10	86.901	65.574	37.880		35.39	C
ATOM	80	NZ	LYS		10	87.937	66.213	38.746	1.00	35.48	N
ATOM	81	N	LYS		11	82.742	68.407	37.167	1.00	33.79	N
MOTA	82	CA	LYS		11	82.841	69.046	35.861		33.06	C
MOTA	83	C	LYS		11	84.280	69.274	35.431	1.00	30.80	C
ATOM	84	0	LYS		11	85.159	69.516	36.259	1.00	30.97	0
ATOM	85	CB	LYS		11	82.072	70.365	35.858	1.00	35.00	C
ATOM	86	CG	LYS		11	80.568	70.163	35.851	1.00		C
ATOM	87	CD	LYS		11	79.802	71.475	35.945	1.00		C
ATOM	88	CE	LYS		11	80.018	72.131	37.301	1.00	47.54	N
ATOM	89	NZ	LYS		11	79.596	71.236 69.168	38.419 34.126	1.00		N
ATOM	90 01	N	GLY		12	84.511	69.370	33.576	1.00		C
ATOM	91	CA C	GLY GLY		12 12	85.837 85.824	70.536	32.604	1.00	26.32	C
ATOM	92 93	0	GLY		12	84.807	70.798	31.952	1.00	25.14	0
ATOM ATOM	93 94	N	TYR		13	86.949	71.241	32.522	1.00		N
	95	CA	TYR		13	87.095	72.389	31.633	1.00		C
ATOM	96	C	TYR		13	88.044	72.084	30.500	1.00	26.35	C
ATOM ATOM	96 97	0	TYR		13	89.042	71.397	30.690	1.00	25.99	0
		CB	TYR		13	87.659	73.604	32.376	1.00		C
MOTA	98 99	CG	TYR		13	86.747	74.148	33.430		34.86	C
ATOM	100		TYR		13	85.471	74.600	33.430		36.83	C
ATOM	101		TYR		13	87.147	74.206	34.763		37.11	C
ATOM	101		TYR		13	84.612	75.093	34.763		39.35	C
ATOM	102		TYR		13	86.294	74.700	35.739		40.44	C
VIOL	103	كتنب	TIK		10	00.274					_

#### **ATOM** TYR A 104 CZ85.027 75.140 13 35.379 1.00 41.53 C **ATOM** 105 TYR A 13 OH 84.161 75.613 36.336 1.00 45.89 0 **ATOM** 106 THR A N 14 87.727 72.606 29.322 1.00 25.75 N **ATOM** THR A 107 88.585 72.448 CA 28.159 1.00 25.34 14 C **ATOM** THR A 108 C 89.024 73.877 27.830 14 1.00 23.61 C MOTA 109 THR A 0 14 88.469 74.832 28.367 1.00 22.92 0 110 ATOM CB THR A 87.817 71.829 26.970 14 1.00 26.77 C ATOM 111 OG1 THR A 88.700 71.699 25.846 14 1.00 31.58 0 **ATOM** THR A 86.635 72.702 112 CG2 14 26.583 1.00 27.27 C **ATOM** 113 PHE A 90.011 15 74.032 N 26.957 1.00 22.10 N **ATOM** PHE A 114 CA 15 90.509 26.613 1.00 20.64 75.353 C MOTA PHE A 15 115 C 89.435 76.391 26.278 1.00 20.42 C **ATOM** PHE A 15 89.579 116 77.555 26.641 0 1.00 20.85 0 PHE A **ATOM** 15 117 CB 91.504 75.238 25.460 1.00 20.90 C **ATOM** 118 15 PHE A 92.685 74.365 CG 25.770 1.00 21.02 C ATOM 119 CD1 PHE A 15 93.568 74.699 26.790 1.00 20.82 C **ATOM** 120 CD2 PHE A 92.897 73.189 15 25.070 1.00 20.78 C **ATOM** 121 CE1 PHE A 15 94.636 73.871 27.102 1.00 20.57 C PHE A **ATOM** 122 15 CE2 93.967 72.356 25.381 C 1.00 20.90 **ATOM** 123 CZPHE A 15 94.832 72.699 26.396 1.00 19.69 $\mathbf{C}$ **ATOM** 124 ASP A 88.362 75.987 25.601 N 16 1.00 20.30 N **ATOM** 125 ASP A 16 87.318 76.936 25.230 CA 1.00 21.26 C **ATOM** 126 ASP A C 26.320 16 86.312 77.314 C 1.00 20.86 **ATOM** 127 ASP A 16 0 85.364 78.050 26.061 0 1.00 20.85 **ATOM** 128 ASP A 76.457 23.983 CB 16 86.564 C 1.00 23.23 1.00 28.01 **ATOM** 129 CG ASP A 16 87.418 76.509 22.721 C **ATOM** OD1 ASP A 88.319 130 22.636 1.00 29.29 16 77.374 0 **ATOM** 131 OD2 ASP A 16 87.167 75.713 21.789 1.00 31.34 0 MOTA 132 ASP A 86.511 76.824 N 17 27.538 1.00 21.11 N ASP A 133 17 85.609 **ATOM** CA 77.165 28.645 1.00 20.51 C ASP A **ATOM** 134 C 17 86.192 78.287 29.487 C 1.00 19.73 1.00 20.34 MOTA 135 ASP A 85.475 78.926 30.244 0 17 O C **ATOM** 136 ASP A 85.376 75.971 29.581 1.00 20.06 17 CB C ASP A 28.913 1.00 21.90 MOTA 137 CG 17 84.651 74.827 OD1 ASP A 83.571 MOTA 28.330 1.00 21.40 0 138 17 75.067 MOTA 139 OD2 ASP A 85.151 28.985 1.00 21.34 0 17 73.680 MOTA VAL A 29.343 140 18 87.491 78.530 1.00 19.47 N N VAL A 79.532 30.149 C 141 88.180 1.00 19.26 ATOM CA 18 C VAL A 80.548 29.410 MOTA 142 C 18 89.054 1.00 20.62 80.344 VAL A 89.468 0 143 28.267 1.00 20.73 MOTA 0 18 C VAL A 89.076 78.833 31.199 ATOM 1.00 18.24 144 СB 18 C CG1 VAL A 88.244 32.060 MOTA 145 18 77.912 1.00 17.13 C 1.00 17.00 CG2 VAL A 90.169 MOTA 78.038 30.505 146 18 N LEU A 81.642 30.106 1.00 20.27 MOTA 147 89.335 N 19 C LEU A 90.186 82.723 29.627 148 1.00 21.23 ATOM CA 19 C LEU A 91.101 83.118 30.785 1.00 21.23 149 **MOTA** C 19 0 LEU A 19 90.700 83.040 31.946 1.00 21.35 **ATOM** 150

ATOM	151	CB	LEU	Α	19	89.347	83.945	29.245	1.00	20.79	C
ATOM	152	CG	LEU	A	19	88.497	83.867	27.991	1.00	20.91	C
ATOM	153	CD1	LEU	A	19	87.558	85.052	27.924	1.00	19.17	C
MOTA	154	CD2	LEU	A	19	89.417	83.812	26.798	1.00	19.74	С
ATOM	155	N	LEU	A	20	92.324	83.537	30.484	1.00	20.61	N
ATOM	156	CA	LEU	Α	20	93.222	83.979	31.541	1.00	21.37	C
ATOM	157	C	LEU	A	20	92.857	85.421	31.868	1.00	22.07	C.
ATOM	158	0	LEU	Α	20	92.623	86.234	30.972	1.00	22.23	0
ATOM	159	CB	LEU	Α	20	94.676	83.894	31.087	1.00	20.78	С
ATOM	160	CG	LEU	Α	20	95.210	82.474	30.972	1.00	21.20	С
ATOM	161	CD1	LEU	A	20	96.567	82.488	30.305	1.00	22.29	C
ATOM	162	CD2	LEU	A	20	95.273	81.857	32.355	1.00	20.57	C
ATOM	163	N	ILE	A	21	92.796	85.734	33.154	1.00	22.61	N
ATOM	164	CA	ILE	A	21	92.442	87.075	33.591	1.00	21.81	C
MOTA	165	C	ILE	Α	21	93.657	87.998	33.597	1.00	23.02	С
ATOM	166	0	ILE	Α	21	94.720	87.649	34.121	1.00	24.26	0
ATOM	167	CB	ILE	Α	21	91.838	87.041	35.010	1.00	21.50	C
ATOM	168	CG1	ILE	A	21	90.667	86.055	35.059	1.00	20.85	C
ATOM	169	CG2	ILE	A	21	91.363	88.434	35.403	1.00	22.33	C
ATOM	170	CD1	ILE	A	21	90.045	85.907	36.432	1.00	19.20	C
ATOM	171	N	PRO	A	22	93.522	89.195	33.010	1.00	23.71	N
ATOM	172	CA	PRO	A	22	94.637	90.141	32.979	1.00	23.92	C
ATOM	173	C	PRO	A	22	95.029	90.494	34.416	1.00	24.84	C
ATOM	174	0	PRO	A	22	94.176	90.501	35.300	1.00	26.16	0
ATOM	175	CB	PRO	A	22	94.033	91.330	32.240	1.00	23.95	C
ATOM	176	CG	PRO	A	22	92.999	90.678	31.355	1.00	24.02	C
ATOM	177	CD	PRO	Α	22	92.346	89.784	32.351	1.00	23.20	C
ATOM	178	N	ALA	A	23	96.308	90.784	34.652	1.00	25.54	N
ATOM	179	CA	ALA	A	23	96.780	91.131	35.995	1.00	25.69	C
ATOM	180	С	ALA		23	97.898	92.164	35.919		26.22	C
ATOM	181	0	ALA		23	98.405	92.446	34.840		25.70	0
ATOM	182	CB	ALA		23	97.273	89.881	36.711	1.00	25.54	C
ATOM	183	N	GLU		24	98.283	92.737	37.055	1.00		N
ATOM	184	CA	GLU		24	99.352	93.725	37.024			C
ATOM	185	С	GLU		24	100.569	93.024	36.449		30.88	С
ATOM	186	0	GLU		24	100.818	91.859	36.753	1.00		0
ATOM	187	CB	GLU		24	99.658	94.276	38.425	1.00	33.75	С
ATOM	188	CG	GLU		24	100.218	93.274	39.406	1.00	39.88	С
ATOM	189	CD	GLU		24	100.497	93.886	40.774		43.61	С
ATOM	190	OE1	GLU		24	101.322	94.825	40.863	1.00		0
ATOM	191	OE2	GLU		24	99.887	93.422	41.763	1.00		0
ATOM	192	N	SER		25	101.315	93.729	35.606	1.00	32.20	N
ATOM	193	CA	SER		25	102.494	93.160	34.971		33.78	C
ATOM	194	C	SER		25	103.714	94.082	35.000		35.09	C
ATOM	195	0	SER		25	103.594	95.289	34.783		35.62	0
ATOM	196	CB	SER		25	102.163	92.803	33.520		32.84	C
ATOM	197	OG	SER	Α	25	103.294	92.273	32.853	1.00	32.95	0

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	MOTA	198	N	HIS	A	26	104.883	93.501	35.267	1.00	36.35	N
	ATOM	199	CA	HIS	A	26	106.136	94.252	35.293	1.00	37.85	C
1	MOTA	200	C	HIS	A	26	107.125	93.640	34.315	1.00	37.72	C
	MOTA	201	0	HIS	A	26	108.333	93.814	34.457	1.00	38.42	0
2	MOTA	202	CB	HIS	Α	26	106.758	94.245	36.685	1.00	40.84	C
2	MOTA	203	CG	HIS	Α	26	105.885	94.852	37.738	1.00	45.91	C
2	MOTA	204	ND1	HIS	Α	26	104.734	94.242	38.195	1.00	48.22	N
2	MOTA	205	CD2	HIS	Α	26	105.978	96.030	38.403	1.00	47.24	C
1	MOTA	206	CE1	HIS	Α	26	104.158	95.017	39.097	1.00	48.99	C
2	MOTA	207	NE2	HIS	A	26	104.892	96.107	39.241	1.00	48.85	N
4	ATOM	208	N	VAL	A	27	106.611	92.922	33.323	1.00	36.55	N
1	MOTA	209	CA	VAL	Α	27	107.462	92.284	32.333	1.00	35.90	C
1	MOTA	210	C	LAV	Α	27	106.851	92.414	30.947	1.00	34.96	C
1	MOTA	211	0	VAL	Α	27	105.689	92.080	30.742	1.00	35.93	0
2	MOTA	212	CB	VAL	A	27	107.665	90.783	32.670	1.00	37.27	C
4	MOTA	213	CG1	JAV	A	27	106.315	90.069	32.733	1.00	38.23	C
	MOTA	214	CG2	LAV	Α	27	108.573	90.130	31.633	1.00	37.72	C
1	MOTA	215	N	LEU	Α	28	107.627	92.922	29.998		33.88	N
1	MOTA	216	CA	LEU	A	28	107.132	93.073	28.638		33.33	C
	MOTA	217	C		A	28	107.190	91.714	27.960		33.40	C
4	MOTA	218	0	LEU		28	107.946	90.842	28.380		33.65	0
2	MOTA	219	CB	LEU		28	107.973	94.093	27.868		32.03	C
	MOTA	220	CG	LEU		28	107.935	95.515	28.434		32.97	C
	MOTA	221				28	108.782	96.432	27.584		31.82	C
	ATOM	222		LEU		28	106.516	96.014	28.468		32.76	C
	MOTA	223	N	PRO		29	106.384	91.508	26.906		33.27	N
	MOTA	224	CA	PRO		29	106.389	90.222	26.211		33.55	C
	MOTA	225	C	PRO		29	107.779	89.774	25.759		34.80	C
	MOTA	226	0	PRO		29	108.142	88.608	25.909		33.98	0
	MOTA	227	CB	PRO		29	105.446	90.480	25.036		33.09	C
	ATOM	228	CG		A	29	104.465	91.448	25.633		32.91	C
	ATOM	229	CD	PRO		29	105.435	92.423	26.248		33.11	C
	ATOM	230	N	ASN		30	108.554	90.712	25.222		36.70	N
	ATOM	231	CA	ASN		30	109.886	90.412	24.713		38.86	C
	ATOM	232	C	ASN		30	110.908	90.077	25.786		39.09	C
	MOTA	233	O	ASN		30	112.034	89.703	25.468		40.70	0
	ATOM	234	CB	ASN		30	110.405	91.574	23.869		40.69	C
	ATOM	235	CG	ASN		30	110.444	92.872	24.644		43.86	0
	MOTA	236				30	109.404	93.408	25.029		46.68	N
	MOTA	237		ASN		30	111.644	93.379	24.892		44.75	N
	ATOM	238	N	GLU		31	110.539	90.205	27.052		38.38 38.11	C
	ATOM	239	CA	GLU		31	111.483	89.876 88.724	28.108 28.984		36.56	C
	ATOM	240	C	GLU		31	111.012		30.028		36.28	0
	ATOM	241	O	GLU		31	111.606 111.784	88.455 91.110	28.966		41.04	C
	ATOM ATOM	242	CB CG	GLU		31 31	111.784	91.828	29.480		44.82	C
	ATOM ATOM	243					110.561		30.265		47.02	C
4	ATOM	244	CD	GLU	A	31	110.307	93.074	30.203	1.00	T1.V4	_

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ATOM	245	OE1	GLU	Α	31	111.466	92.948	31.376	1.00 49.39	0
MOTA	246	OE2	GLU	Α	31	110.629	94.184	29.765	1.00 48.56	0
MOTA	247	N	VAL	Α	32	109.942	88.045	28.572	1.00 34.46	N
ATOM	248	CA	VAL	Α	32	109.469	86.912	29.354	1.00 32.34	C
ATOM	249	C	VAL	A	32	110.435	85.761	29.082	1.00 32.04	C
ATOM	250	0	VAL	A	32	110.987	85.648	27.978	1.00 30.55	0
ATOM	251	CB	VAL	Α	32	108.040	86.463	28.957	1.00 31.69	C
ATOM	252	CG1	VAL	Α	32	107.056	87.578	29.170	1.00 31.19	C
ATOM	253	CG2	VAL	Α	32	108.023	86.034	27.535	1.00 34.01	C
ATOM	254	N	ASP	A	33	110.642	84.918	30.092	1.00 31.28	N
ATOM	255	CA	ASP	Α	33	111.535	83.773	29.973	1.00 31.91	C
ATOM	256	C	ASP	Α	33	110.738	82.549	29.523	1.00 31.62	C
ATOM	257	0	ASP	Α	33	109.861	82.079	30.247	1.00 32.49	0
ATOM	258	CB	ASP	A	33	112.191	83.491	31.326	1.00 32.88	C
ATOM	259	CG	ASP	Α	33	113.233	82.397	31.252	1.00 33.65	C
ATOM	260	OD1	ASP	A	33	113.723	81.983	32.322	1.00 35.11	0
MOTA	261	OD2	ASP	Α	33	113.569	81.959	30.128	1.00 34.94	0
ATOM	262	N	LEU	Α	34	111.038	82.037	28.332	1.00 31.40	N
ATOM	263	CA	LEU	Α	34	110.334	80.871	27.806	1.00 31.53	C
ATOM	264	С	LEU	Α	34	110.977	79.528	28.149	1.00 33.19	С
ATOM	265	0	LEU	A	34	110.436	78.470	27.801	1.00 33.53	0
ATOM	266	CB	LEU	A	34	110.194	80.979	26.285	1.00 30.67	C
ATOM	267	CG	LEU	Α	34	108.990	81.696	25.671	1.00 29.76	С
ATOM	268	CD1	LEU	A	34	108.758	83.013	26.322	1.00 31.73	C
ATOM	269	CD2	LEU	Α	34	109.229	81.865	24.192	1.00 29.76	C
ATOM	270	N	LYS	A	35	112.122	79.550	28.826	1.00 33.83	N
ATOM	271	CA	LYS	Α	35	112.780	78.291	29.159	1.00 34.29	C
MOTA	272	С	LYS		35	111.974	77.434	30.114	1.00 33.40	C
ATOM	273	0	LYS		35	111.196	77.941	30.934	1.00 32.21	0
ATOM	274	CB	LYS		35	114.167	78.521	29.770	1.00 36.57	C
ATOM	275	CG	LYS		35	115.196	79.107	28.821	1.00 39.86	C
ATOM	276	CD	LYS		35	116.586	79.017	29.440	1.00 43.11	C
ATOM	277	CE	LYS		35	116.664	79.750	30.780	1.00 45.95	C
ATOM	278	NZ	LYS		35	116.444	81.232	30.640	1.00 49.28	N
ATOM	279	N	THR		36	112.169	76.124	30.005	1.00 31.86	N
ATOM	280	CA	THR		36	111.484	75.190	30.886	1.00 30.97	C
ATOM	281	C	THR		36	112.381	73.969	31.106	1.00 31.69	C
MOTA	282	0	THR		36	112.895	73.385	30.146	1.00 31.07	0
ATOM	283	CB	THR		36	110.092	74.754	30.298	1.00 28.79	C
ATOM	284	OG1			36	109.436	73.879	31.223	1.00 26.65	0
MOTA	285	CG2	THR		36	110.257	74.036	28.956	1.00 25.84	C
ATOM	286	N	LYS		37	112.603	73.617	32.368	1.00 32.29	N
ATOM	287	CA	LYS		37	113.418	72.450	32.675	1.00 34.79	C
ATOM	288	C	LYS		37	112.456	71.305	32.920	1.00 34.78	0
ATOM	289	0	LYS		37	111.611	71.375	33.812	1.00 35.62	C
ATOM	290	CB	LYS		37	114.291	72.670	33.921	1.00 36.21	G
ATOM	291	CG	LYS	А	37	115.164	71.452	34.243	1.00 39.26	Ç

ATOM	292	CD	LYS	A	37	116.208	71.705	35.329	1.00	41.30	C
ATOM	293	CE	LYS	A	37	115.590	71.952	36.693	1.00	43.41	C
ATOM	294	NZ	LYS	Α	37	116.654	72.208	37.723	1.00	45.01	N
ATOM	295	N	LEU	Α	38	112.575	70.261	32.109	1.00	35.03	N
ATOM	296	CA	LEU	Α	38	111.702	69.102	32.219	1.00	35.59	C
ATOM	297	C	LEU	A	38	112.319	68.043	33.115	1.00	36.52	C
ATOM	298	0	LEU	Α	38	111.614	67.298	33.796	1.00	36.39	0
ATOM	299	CB	LEU	Α	38	111.423	68.533	30.825	1.00	33.92	С
ATOM	300	CG	LEU	Α	38	110.683	69.502	29.893	1.00	32.62	С
MOTA	301	CD1	LEU	A	38	110.462	68.861	28.531	1.00	31.57	C
ATOM	302	CD2	LEU	A	38	109.356	69.887	30.523	1.00	31.42	C
ATOM	303	N	ALA	A	39	113.644	67.995	33.106	1.00	38.25	N
ATOM	304	CA	ALA	A	39	114.403	67.056	33.915	1.00	40.08	C
ATOM	305	C	ALA	A	39	115.831	67.591	34.003	1.00	41.95	C
ATOM	306	0	ALA	A	39	116.223	68.463	33.220	1.00	42.09	0
ATOM	307	CB	ALA	A	39	114.388	65.677	33.266	1.00	39.28	C
ATOM	308	N	ASP	A	40	116.602	67.073	34.954	1.00	43.87	N
ATOM	309	CA	ASP	A	40	117.993	67.486	35.146	1.00	45.01	C
ATOM	310	С	ASP	A	40	118.722	67.562	33.806	1.00	45.03	C
ATOM	311	0	ASP	A	40	119.467	68.505	33.535	1.00	45.51	0
ATOM	312	CB <sub>.</sub>	ASP	A	40	118.688	66.481	36.060	1.00	46.91	C
MOTA	313	CG	ASP	A	40	118.075	66.436	37.446	1.00	49.68	C
ATOM	314	OD1	ASP	A	40	118.340	65.458	38.180	1.00	51.95	0
ATOM	315	OD2	ASP	Α	40	117.345	67.387	37.812	1.00	50.80	0
MOTA	316	N	ASN	A	41	118.487	66.557	32.972	1.00	44.94	N
ATOM	317	CA	ASN	A	41	119.090	66.455	31.642	1.00	44.54	C
ATOM	318	С	ASN		41	118.289	67.153	30.545	1.00		C
ATOM	319	0	ASN		41	118.775	67.316	29.422		42.49	0
MOTA	320	CB	ASN		41	119.233	64.969	31.280		47.24	C
ATOM	321	CG	ASN		41	118.956	64.687	29.801		48.33	C
MOTA	322	OD1	ASN		41	119.714	65.095	28.922	1.00		0
ATOM	323	ND2	ASN		41	117.852	64.001	29.530		47.16	N
MOTA	324	N	LEU		42	117.074	67.589	30.866		41.33	N
ATOM	325	CA	LEU		42	116.227	68.186	29.845		39.00	C
ATOM	326	С	LEU		42	115.637	69.573	30.120	1.00		C
MOTA	327	0	LEU		42	114.653	69.716	30.848		37.43	0
MOTA	328	CB	LEU		42	115.102	67.200	29.532		38.42	C
ATOM	329	CG	LEU		42	114.599	67.104	28.099	1.00		C
ATOM	330	CD1	LEU		42	115.752	66.736	27.172		36.38	C
ATOM	331	CD2	LEU		42	113.509	66.047	28.032		36.67	C
ATOM	332	N	THR		43	116.242	70.586	29.511		35.65	N
ATOM	333	CA	THR		43	115.783	71.960	29.639		34.45	C
MOTA	334	C	THR		43	115.610	72.521	28.232		33.69	C
ATOM	335	0	THR		43	116.539	72.482	27.421		33.77	0
ATOM	336	CB	THR		43	116.792	72.827	30.404		34.76	C
ATOM	337	OG1	THR		43	116.874	72.372	31.755		36.40	0
MOTA	338	CG2	THR	Α	43	116.358	74.281	30.402	1.00	35.16	C

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	MOTA	339	N	LEU	A	44	114.421	73.039	27.936	1.00	31.23	N
	MOTA	340	CA	LEU	Α	44	114.157	73.578	26.614	1.00	28.88	C
	MOTA	341	C	LEU	A	44	114.097	75.095	26.683	1.00	27.97	C
	MOTA	342	0	LEU	A	44	113.799	75.655	27.733	1.00	29.11	0
	ATOM	343	CB	LEU	A	44	112.830	73.016	26.094	1.00	28.32	C
	ATOM	344	CG	LEU	A	44	112.680	71.487	26.107	1.00	27.65	C
	ATOM	345	CD1	LEU	A	44	111.299	71.107	25.598	1.00	26.79	C
	MOTA	346	CD2	LEU	Α	44	113.756	70.840	25.248	1.00	28.11	C
	ATOM	347	N	ASN	A	45	114.380	75.756	25.565	1.00	26.86	N
	ATOM	348	CA	ASN	Α	45	114.337	77.215	25.492	1.00	27.18	C
	ATOM	349	C	ASN	A	45	112.933	77.752	25.191	1.00	27.96	C
	ATOM	350	0	ASN	A	45	112.680	78.947	25.312	1.00	27.92	0
	ATOM	351	CB	ASN	Α	45	115.323	77.699	24.443	1.00	27.75	C
	ATOM	352	CG	ASN	A	45	116.755	77.510	24.878	1.00	28.46	C
	ATOM	353	OD1	ASN	A	45	117.644	77.325	24.051	1.00	30.13	0
	ATOM	354	ND2	ASN		45	116.993	77.579	26.184	1.00	27.73	N
	ATOM	355	N	ILE		46	112.033	76.863	24.777	1.00	27.88	N
	ATOM	356	CA	ILE		46	110.636	77.213	24.516	1.00	26.86	C
	ATOM	357	C			46	109.848	75.975	24.946	1.00	27.09	C
	ATOM	358	0	ILE		46	110.282	74.843	24.701	1.00	27.35	0
	ATOM	359	СВ	ILE		46	110.375	77.537	23.028	1.00	25.10	С
	ATOM	360	CG1			46	110.724	76.340	22.147	1.00	25.03	C
	ATOM	361	CG2	ILE		46	111.182	78.756	22.632	1.00	26.47	C
	ATOM	362	CD1	ILE	A	46	110.468	76.573	20.657	1.00		С
	ATOM	363	N	PRO		47	108.702	76.170	25.626		25.80	N
	ATOM	364	CA	PRO		47	107.849	75.082	26.112	1.00	24.42	C
•	ATOM	365	C	PRO		47	107.077	74.339	25.024	1.00	24.61	С
	ATOM	366	0	PRO		47	105.912	73.988	25.223	1.00	23.97	0
	ATOM	367	СВ	PRO		47	106.929	75.808	27.083	1.00	23.99	C
	ATOM	368	CG	PRO		47	106.697	77.097	26.351	1.00	23.63	C
	ATOM	369	CD	PRO		47	108.135	77.467	26.036	1.00	24.98	C
	ATOM	370	N			48	107.735	74.063	23.901	1.00	23.91	N
	ATOM	371	CA	ILE		48	107.083	73.395	22.784	1.00	24.05	C
	ATOM	372	C			48	107.730	72.083	22.322	1.00	24.29	C
	ATOM	373	0			48	108.943	72.013	22.098	1.00	23.94	0
	ATOM	374	СВ	ILE		48	106.997	74.368	21.592	1.00	25.68	С
	ATOM	375	CG1	ILE		48	106.193	75.605	22.009	1.00	25.65	С
	ATOM	376	CG2			48	106.376	73.675	20.379	1.00	25.11	C
	ATOM	377	CD1	ILE		48	106.124	76.678	20.949	1.00	26.76	C
	ATOM	378	N	ILE	A	49	106.897	71.054	22.162	1.00	23.20	N
	ATOM	379	CA	ILE		49	107.343	69.732	21.725	1.00	23.13	С
	ATOM	380	C	ILE		49	106.454	69.215	20.588	1.00	23.01	С
	ATOM	381	0	ILE		49	105.241	69.397	20.623		22.62	0
	ATOM	382	CB	ILE		49	107.274	68.723	22.876		22.04	C
	ATOM	383	CG1			49	108.027	69.277	24.093		20.85	C
	ATOM	384		ILE		49	107.847	67.376	22.412		21.51	C
	ATOM	385		ILE		49	107.922	68.416	25.333		19.42	C
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#### 68.573 **ATOM** 386 N THR A 50 107.059 19.587 1.00 22.84 N 387 THR A 50 106.303 68.049 **ATOM** C CA 18.449 1.00 23.78 THR A C **ATOM** 388 C 50 105.913 66.582 18.643 1.00 24.70 THR A **ATOM** 389 65.746 1.00 25.64 0 50 106.746 18.989 0 THR A 107.096 **ATOM** 390 CB 50 68.200 17.126 1.00 23.25 C THR A 391 50 108.344 67.503 17.219 1.00 24.27 **ATOM** OG1 0 69.666 1.00 23.48 THR A **ATOM** 392 CG2 50 107.360 16.838 C 104.641 **ATOM** 393 51 66.279 18.401 1.00 24.84 ALA A N N **ATOM** 394 CA ALA A 51 104.100 64.937 18.576 1.00 24.13 C ALA A 63.813 C **ATOM** 395 C 51 104.842 17.849 1.00 25.28 63.997 0 ALA A 51 105.385 16.760 1.00 24.45 0 ATOM 396 ALA A 397 51 102.638 C 64.931 18.183 1.00 23.17 ATOM CB MOTA ALA A 52 104.852 62.639 18.477 1.00 26.02 N 398 Ŋ 105.496 ALA A 52 1.00 26.54 C **ATOM** 399 CA 61.450 17.932 400 52 104.539 60.787 16.945 1.00 26.47 C **ATOM** C ALA A 401 ALA A 52 104.097 59.660 17.146 1.00 26.78 0 MOTA 0 ALA A 52 105.833 1.00 25.79 C 402 CB 60.484 19.063 MOTA **HETATM** 403 53 61.496 N N MSE A 104.221 15.871 1.00 27.09 MSE A 53 103.294 60.985 14.868 1.00 27.91 C **HETATM** 404 CA MSE A 103.997 60.874 1.00 27.12 C **HETATM** 405 C 53 13.532 MSE A 53 104.710 1.00 26.01 0 0 **HETATM** 406 61.790 13.133 MSE A 1.00 29.38 53 61.932 C **HETATM** 407 CB 102.094 14.775 MSE A 1.00 32.60 C 408 CG 53 101.381 62.108 16.124 **HETATM** SE SE MSE A 53 99.997 16.089 1.00 36.50 **HETATM** 409 63.279 100.826 C CE MSE A 53 64.714 15.572 1.00 38.59 **HETATM** 410 59.757 ASP A 54 103.790 12.837 1.00 27.85 411 N N MOTA 412 ASP A 1.00 28.83 C **ATOM** 104.443 59.546 11.550 CA 54 10.460 ASP A 54 104.108 60.560 1.00 29.48 C **ATOM** 413 C 54 9.366 **ATOM** 0 ASP A 104.665 60.493 1.00 30.74 414 ASP A 104.212 1.00 29.72 11.049 C MOTA 415 CB 54 58.113 10.826 ASP A MOTA 102.754 57.784 416 CG 54 1.00 31.41 56.604 10.539 OD1 ASP A 102.474 1.00 34.67 MOTA 417 54 101.888 OD2 ASP A 58.674 1.00 32.27 0 ATOM 418 10.925 54 10.753 THR A 55 103.211 61.501 1.00 29.32 N ATOM 419 N 1.00 27.93 THR A 55 102.850 62,548 9.796 C 420 CA MOTA 103.282 C MOTA C THR A 55 63.914 10.328 1.00 27.76 421 9.728 1.00 27.97 422 102.985 0 MOTA 0 THR A 55 64.945 423 THR A 62.574 9.497 C 55 101.328 1.00 28.66 MOTA ÇВ 100.584 OG1 THR A 55 62.643 10.721 0 MOTA 424 1.00 28.50 8.720 CG2 THR A 55 100.915 61.334 C MOTA 425 1.00 28.74 VAL A 426 N MOTA N 56 103.997 63.912 11.454 1.00 26.53 12.061 1.00 25.82 C **ATOM** 427 VAL A 65.153 56 104.479 CA C VAL A MOTA 428 C 56 105.975 65.173 12.392 1.00 25.65 VAL A 0 429 1.00 27.06 0 56 106.675 66.085 11.977 MOTA C MOTA 430 CBVAL A 56 103.700 65.500 13.361 1.00 24.87 C CG1 VAL A 104.243 66.781 13.975 1.00 23.59 **MOTA** 431 56 C 102.230 65.679 ATOM 432 CG2 VAL A 56 13.052 1.00 24.94

1.00 26.14

#### 64.178 1.00 25.55 THR A 106.487 13.108 N 433 N 57 ATOM CA THR A 57 107.895 64.235 13.479 1.00 24.90 C **MOTA** 434 1.00 25.70 C 435 THR A **ATOM** C 57 108.881 63.190 12.993 **ATOM** 0 THR A 57 108.933 62.077 13.510 1.00 26.76 0 436 THR A 57 108.070 64.304 15.022 C **ATOM** 1.00 24.37 437 CB OG1 THR A 0 **ATOM** 57 107.342 65.420 15.536 1.00 24.74 438 109.535 THR A 64.499 15.391 1.00 22.67 C 439 57 **ATOM** CG2 109.696 63.592 N **ATOM** 58 12.025 1.00 25.19 440 N GLY A 110.762 C 441 GLY A 58 62.749 1.00 24.08 **ATOM** CA 11.517 GLY A C 112.004 63.541 **ATOM** 11.912 1.00 23.78 442 C 58 12.682 1.00 21.95 0 **ATOM** 443 0 GLY A 58 111.881 64.497 113.180 1.00 24.38 SER A 63.179 N **ATOM** 11.399 444 N 59 11.723 C SER A 59 114.417 63.901 1.00 24.47 **ATOM** 445 CA 114.406 1.00 24.98 C 65.388 11.318 446 SER A 59 ATOM C 66.229 12.052 0 **ATOM** 0 SER A 59 114.915 1.00 25.40 447 1.00 23.93 SER A 59 115.614 63.193 C 11.090 MOTA 448 CB 9.705 1.00 26.47 SER A 59 115.404 62.980 0 MOTA 449 OG 10.156 1.00 25.85 N LYS A 60 **ATOM** 113.836 65.710 450 N 9.702 1.00 26.54 C LYS A 113.747 67.103 CA 60 ATOM 451 10.696 112.985 67.987 1.00 25.73 C 452 LYS A C 60 MOTA LYS A 69.118 10.983 1.00 25.68 0 60 113.389 MOTA 453 0 C LYS A 60 8.358 1.00 27.44 MOTA 113.029 67.190 454 CB 7.148 1.00 31.91 C 455 LYS A 113.902 67.124 60 **ATOM** CG 5.878 1.00 35.71 C CD LYS A 113.034 67.048 ATOM 456 60 C 1.00 36.50 LYS A 68.237 5.710 MOTA 60 112.071 457 CE И LYS A 60 1.00 38.79 **ATOM** 112.742 69.542 5.417 458 NZ N 459 MSE A 111.864 67.482 11.196 1.00 25.26 N 61 **HETATM** 1.00 25.61 C MSE A 111.060 68.242 **HETATM** 12.140 460 CA 61 111.773 68.317 13.482 1.00 24.55 MSE A **HETATM** 61 461 C 1.00 24.99 0 MSE A 61 111.805 14.115 69.373 **HETATM** 462 0 MSE A 109.678 67.587 12.295 1.00 27.32 **HETATM** 463 CB 61 1.00 29.47 68.290 13.258 C MSE A 61 108.708 **HETATM** 464 CG 12.808 1.00 35.03 SE 108.189 69.988 **HETATM** MSE A 61 465 SE C MSE A 70.965 13.376 1.00 35.39 466 61 109.528 HETATM CE И 112.343 67.194 13.914 1.00 23.85 ALA A 62 MOTA 467 $\mathbf{N}$ C 1.00 23.20 ALA A 113.065 468 62 67.144 15.184 ATOM CA C 114.223 68.124 15.141 1.00 22.77 ATOM ALA A 62 469 C 1.00 22.10 114.588 16.147 ATOM 470 ALA A 62 68.717 0 C 113.581 1.00 23.29 471 ALA A 65.747 15.445 CB 62 ATOM 1.00 22.99 N 68.292 ATOM ILE A 63 114.798 13.959 472 N 1.00 23.52 C 115.898 473 69.221 13.775 MOTA CA ILE A 63 C 70.683 13.828 474 ILE A 63 115.426 1.00 24.19 ATOM C 0 71.519 14.514 ILE A 116.042 1.00 23.98 475 63 MOTA 0 C ILE A 63 116.606 68.934 12.442 1.00 23.78 MOTA 476 CB C 12.569 1.00 24.18 CG1 ILE A 63 117.400 67.627 ATOM 477 C CG2 ILE A 63 117.476 70.110 12.047 1.00 25.35 MOTA 478

118.130

CD1 ILE A 63

ATOM

479

67.209

11.319

ATOM	480	N	ALA	A	64	114.326	70.981	13.129	1.00 24.07	N
ATOM	481	CA	ALA	A	64	113.762	72.335	13.091	1.00 23.68	С
ATOM	482	C	ALA	A	64	113.230	72.799	14.447	1.00 23.69	С
ATOM	483	0	ALA	A	64	113.452	73.943	14.847	1.00 24.51	0
ATOM	484	CB	ALA	A	64	112.649	72.418	12.052	1.00 22.06	C
ATOM	485	N	ILE	A	65	112.528	71.927	15.161	1.00 23.23	N
MOTA	486	CA	ILE	A	65	111.993	72.336	16.456	1.00 23.60	C
ATOM	487	C	ILE	A	65	113.100	72.513	17.500	1.00 23.66	C
ATOM	488	0	ILE	A	65	112.983	73.332	18.416	1.00 23.86	0
ATOM	489	CB	ILE	A	65	110.932	71.324	16.980	1.00 23.74	С
ATOM	490	CG1	ILE	A	65	110.372	71.801	18.326	1.00 23.72	C
ATOM	491	CG2	ILE	A	65	111.541	69.929	17.108	1.00 23.00	С
ATOM	492	CD1	ILE	A	65	109.618	73.128	18.258	1.00 23.28	C
ATOM	493	N	ALA	A	66	114.169	71.739	17.370	1.00 23.35	N
ATOM	494	CA	ALA	A	66	115.277	71.845	18.310	1.00 24.29	C
ATOM	495	C	ALA	A	66	116.019	73.163	18.037	1.00 23.90	С
ATOM	496	0	ALA		66	116.379	73.888	18.970	1.00 23.78	0
ATOM	497	CB	ALA		66	116.222	70.639	18.164	1.00 24.32	С
ATOM	498	N	ARG		67	116.236	73.480	16.762	1.00 22.77	N
ATOM	499	CA	ARG		67	116.916	74.725	16.415	1.00 23.18	C
ATOM	500	C	ARG		67	116.158	75.926	16.954	1.00 23.38	C
ATOM	501	0	ARG		67	116.747	76.955	17.274	1.00 23.27	0
ATOM	502	CB	ARG		67	117.061	74.850	14.900	1.00 23.27	C
ATOM	503	CG	ARG		67	118.069	73.896	14.321	1.00 25.02	C
ATOM	504	CD	ARG		67	118.110	73.962	12.806	1.00 25.68	C
ATOM	505	NE	ARG		67	119.304	73.292	12.310	1.00 27.14	N
ATOM	506	CZ	ARG		67	119.562	73.049	11.032	1.00 28.72	C
ATOM	507	NH1	ARG		67	118.707	73.415	10.088	1.00 30.38	N
ATOM	508	NH2	ARG		67	120.694	72.451	10.698	1.00 30.93	N
ATOM	509	N	ALA		68	114.842	75.791	17.053	1.00 23.80	N
ATOM	510	CA	ALA		68	114.006	76.871	17.555	1.00 24.57	C
ATOM	511	C	ALA		68	114.011	76.902	19.083 19.691	1.00 24.29 1.00 25.66	0
ATOM	512	O	ALA		68 68	113.504 112.582	77.839 76.714	17.032	1.00 25.88	C
ATOM	513	CB N	ALA GLY		69	114.582	75.876	19.704	1.00 24.33	N
ATOM ATOM	514 515	CA	GLY		69	114.582	75.844	21.154	1.00 24.27	C
ATOM	516	C	GLY		69	113.718	74.804	21.764	1.00 23.20	C
ATOM	517	0	GLY		69	113.710	74.691	22.986	1.00 23.02	0
ATOM	518	N	GLY		70	113.033	74.031	20.918	1.00 23.60	N
ATOM	519	CA	GLY		70	112.124	73.025	21.416	1.00 23.82	C
ATOM	520	C	GLY		70	112.661	71.607	21.404	1.00 24.81	C
ATOM	521	o	GLY		70	113.873	71.383	21.370	1.00 24.12	0
ATOM	522	N	LEU		71	111.745	70.643	21.430	1.00 24.57	N
ATOM	523	CA	LEU		71	112.113	69.235	21.436	1.00 24.66	C
ATOM	524	C	LEU		71	111.237	68.454	20.473	1.00 25.01	C
ATOM	525	0	LEU		71	110.023	68.641	20.443	1.00 26.48	0
ATOM	526	CB	LEU		71	111.945	68.653	22.842	1.00 23.51	C
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ATOM	527	CG	LEU	A	71	112.350	67.193	23.025	1.00	23.35	С
ATOM	528	CD1	LEU	Α	71	113.832	67.082	22.735	1.00	24.00	C
ATOM	529	CD2	LEU	Α	71	112.053	66.716	24.441	1.00	21.93	С
MOTA	530	N	GLY	Α	72	111.858	67.589	19.682	1.00	24.71	N
ATOM	531	CA	GLY	A	72	111.110	66.767	18.751	1.00	24.20	C
ATOM	532	C	GLY		72	111.122	65.316	19.216	1.00		C
ATOM	533	0	GLY		72	112.148	64.816	19.671	1.00		0
MOTA	534	N	VAL		73	109.980	64.643	19.116	1.00		N
ATOM	535	CA	LAV		73	109.874	63.247	19.519	1.00	23.30	С
ATOM	536	C	LAV		73	109.683	62.388	18.269	1.00		C
ATOM	537	0	VAL		73	108.610	62.387	17.664	1.00		0
ATOM	538	СВ	VAL		73	108.680	63.043	20.496	1.00		C
ATOM	539	CG1	VAL		73	108.604	61.590	20.950	1.00	22.04	C
ATOM	540	CG2	LAV		73	108.835	63.970	21.703	1.00		C
ATOM	541	N	ILE		74	110.742	61.680	17.874	1.00		N
ATOM	542	CA	ILE		74	110.698	60.822	16.690	1.00		C
ATOM	543	C			74	109.751	59.668	16.939	1.00		C
ATOM	544	0	ILE		74	109.871	58.932	17.917	1.00		0
ATOM	545	CB	ILE		74	112.070	60.216	16.332	1.00		C
MOTA	546	CG1	ILE		74	113.127	61.317	16.167	1.00		C
ATOM	547	CG2	ILE		74	111.934	59.376	15.058	1.00	25.49	C
ATOM	548	CD1	ILE		74	112.841	62.293	15.058		29.69	C
MOTA	549	N	HIS		75	108.808	59.517	16.029	1.00		N
MOTA	550	CA	HIS		75	107.800	58.482	16.111		34.55	C
ATOM	551	C	HIS		75	108.375	57.081	15.924		34.59	C
ATOM	552	0	HIS		75	109.427	56.904	15.301		34.10	0
ATOM	553	CB	HIS		75 75	106.758	58.798	15.065		39.30	C
ATOM	554	CG	HIS		75 75	107.336	59.466	13.861		46.72	C
ATOM	555	ND1	HIS		75 75	108.504	59.030	13.267		50.41	N
ATOM	556	CD2	HIS		75 75	106.850	60.439	13.055		50.89	C
ATOM	557	CE1	HIS		75 75	108.706	59.698	12.143		51.89 53.11	C N
ATOM	558	NE2 N	HIS LYS		75 76	107.716 107.666	60.561 56.086	11.990 16.458		34.75	N
ATOM ATOM	559 560	CA	LYS		76	107.000	54.688	16.409		35.86	C
ATOM	561	C	LYS		76	107.398	53.821	15.349		36.31	C
ATOM	562	0	LYS		76	107.467	52.586	15.391		36.33	0
ATOM	563	CB	LYS		76	107.896	54.063	17.790		35.61	C
ATOM	564	CG	LYS		76	106.439	54.003	18.235		36.48	C
ATOM	565	CD	LYS		76	106.313	53.405	19.624		38.36	C
ATOM	566	CE	LYS		76	104.892	52.950	19.910		38.72	C
ATOM	567	NZ	LYS		76	103.918	54.041	19.706		40.04	N
ATOM	568	N	ASN		77	106.736	54.461	14.394		36.56	N
ATOM	569	CA	ASN		77	106.730	53.723	13.360		36.37	C
ATOM	570	CA	ASN		77	107.032	53.725	12.265		36.27	C
ATOM	571	0	ASN		77	106.823	53.619	11.077		36.44	0
ATOM	572	CB	ASN		77	104.904	54.587	12.784		37.00	C
ATOM	573	CG	ASN		77	103.721	53.770	12.312		37.48	C
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ATOM	574	OD1	ASN	A	77	103.882	52.747	11.647	1.00 38.78	0
ATOM	575	ND2	ASN	Α	77	102.520	54.232	12.634	1.00 37.68	N
HETATM	576	N	MSE	Α	78	108.132	52.730	12.681	1.00 35.58	N
HETATM	577	CA	MSE	Α	78	109.177	52.293	11.758	1.00 36.20	С
HETATM	578	C	MSE	Α	78	110.104	51.313	12.475	1.00 35.86	С
HETATM	579	0	MSE	Α	78	110.038	51.188	13.694	1.00 36.31	0
HETATM	580	CB	MSE	Α	78	109.956	53.500	11.207	1.00 37.66	C
HETATM	581	CG	MSE	Α	78	110.414	54.534	12.237	1.00 39.57	C
<b>HETATM</b>	582	SE	MSE	Α	78	111.321	55.923	11.449	1.00 43.88	SE
<b>HETATM</b>	583	CE	MSE	Α	78	111.298	57.131	12.764	1.00 43.70	С
ATOM	584	N	SER	Α	79	110.956	50.606	11.739	1.00 34.43	N
ATOM	585	CA	SER	Α	79	111.828	49.639	12.391	1.00 34.01	C
ATOM	586	C	SER	Α	79	112.734	50.315	13.421	1.00 34.22	C
ATOM	587	0	SER	Α	79	112.961	51.525	13.356	1.00 34.05	0
ATOM	588	CB	SER	Α	79	112.676	48.897	11.362	1.00 32.01	C
ATOM	589	OG	SER	Α	79	113.613	49.764	10.771	1.00 31.21	0
ATOM	590	N	ILE	Α	80	113.229	49.528	14.378	1.00 33.98	N
ATOM	591	CA	ILE	Α	80	114.123	50.031	15.418	1.00 34.25	C
ATOM	592	С	ILE	Α	80	115.356	50.641	14.759	1.00 34.44	С
ATOM	593	0	ILE	Α	80	115.847	51.687	15.177	1.00 34.28	0
ATOM	594	CB	ILE	Α	80	114.574	48.895	16.369	1.00 33.95	C
ATOM	595	CG1	ILE	A	80	113.366	48.322	17.105	1.00 33.91	C
ATOM	596	CG2	ILE	A	80	115.594	49.416	17.369	1.00 34.09	C
ATOM	597	CD1	ILE	Α	80	113.714	47.217	18.090	1.00 34.10	C
ATOM	598	N	THR	A	81	115.849	49.976	13.720	1.00 34.69	N
ATOM	599	CA	THR	Α	81	117.017	50.450	12.991	1.00 34.33	C
ATOM	600	С	THR	A	81	116.719	51.788	12.306	1.00 34.17	C
ATOM	601	0	THR	A	81	117.533	52.708	12.365	1.00 34.07	0
ATOM	602	CB	THR	Α	81	117.472	49.393	11.944	1.00 33.53	C
ATOM	603	OG1	THR	A	81	117.988	48.248	12.627	1.00 33.84	0
ATOM	604	CG2	THR	A	81	118.542	49.942	11.036	1.00 33.63	C
ATOM	605	N	GLU	Α	82	115.550	51.895	11.674	1.00 34.36	N
ATOM	606	CA	GLU	A	82	115.148	53.117	10.973	1.00 33.72	C
ATOM	607	С	GLU	A	82	114.936	54.269	11.941	1.00 32.06	C
ATOM	608	0	GLU	A	82	115.289	55.407	11.642	1.00 31.19	0
ATOM	609	СВ	GLU	Α	82	113.853	52.884	10.188	1.00 37.28	С
ATOM	610	CG	GLU	Α	82	113.953	51.828	9.093	1.00 42.63	C
ATOM	611	CD	GLU	Α	82	112.589	51.420	8.520	1.00 45.15	C
ATOM	612	OE1	GLU	Α	82	111.711	50.979	9.302	1.00 44.54	0
ATOM	613	OE2	GLU	Α	82	112.404	51.530	7.284	1.00 47.14	0
ATOM	614	N	GLN	Α	83	114.354	53.978	13.100	1.00 30.48	N
ATOM	615	CA	GLN	A	83	114.107	55.022	14.080	1.00 29.17	C
ATOM	616	C	GLN		83	115.413	55.520	14.694	1.00 29.61	С
ATOM	617	0	GLN		83	115.565	56.720	14.947	1.00 30.50	0
ATOM	618	СВ	GLN		83	113.178	54.526	15.186	1.00 27.78	С
ATOM	619	CG	GLN		83	112.691	55.646	16.087	1.00 26.47	С
ATOM	620	CD	GLN		83	111.897	55.160	17.277	1.00 25.24	C

MOTA	621	OE1	GLN	A	83	111.368	55.960	18.048	1.00	27.62	0
MOTA	622	NE2	GLN	A	83	111.817	53.851	17.444	1.00	25.23	N
MOTA	623	N	ALA	Α	84	116.353	54.604	14.929	1.00	29.43	N
ATOM	624	CA	ALA	A	84	117.654	54.955	15.501	1.00	28.41	С
MOTA	625	C	ALA	Α	84	118.472	55.837	14.559	1.00	28.88	C
MOTA	626	0	ALA	A	84	119.146	56.766	14.998	1.00	29.24	0
MOTA	627	CB	ALA	Α	84	118.427	53.704	15.826	1.00	27.73	C
MOTA	628	N	GLU	A	85	118.416	55.535	13.267	1.00	28.73	N
ATOM	629	CA	GLU	A	85	119.139	56.298	12.260	1.00	31.28	C
ATOM	630	C	GLU	A	85	118.518	57.684	12.123	1.00	31.60	C
MOTA	631	0	GLU	A	85	119.203	58.673』	11.854	1.00	30.88	0
ATOM	632	CB	GLU	A	85	119.072	55.557	10.922	1.00	34.79	C
ATOM	633	CG	GLU	A	85	119.767	56.251	9.762	1.00	40.50	C
MOTA	634	CD	GLU	Α	85	121.231	56.553	10.051	1.00	43.61	С
MOTA	635	OE1	GLU	Α	85	121.984	55.612	10.413		44.96	0
MOTA	636	OE2	GLU	A	85	121.624	57.736	9.905		44.87	0
ATOM	637	N	GLU		86	117.204	57.739	12.300		31.99	N
ATOM	638	CA	GLU		86	116.452	58.983	12.232		32.20	C
ATOM	639	С	GLU		86	116.996	59.854	13.382	1.00		С
ATOM	640	0	GLU		86	117.257	61.053	13.218		32.08	0
ATOM	641	CB	GLU		86	114.972	58.667	12.449		33.98	C
MOTA	642	CG	GLU		86	114.002	59.724	11.988		38.81	C
MOTA	643	CD	GLU		86	113.971	59.856	10.480	1.00		C
MOTA	644	OE1	GLU		86	113.133	60.628	9.969	1.00		0
ATOM	645	OE2	GLU	A	86	114.785	59.190	9.807	1.00		0
ATOM	646	N	VAL		87	117.176	59.225	14.543			N
ATOM	647	CA	VAL		87	117.701	59.890	15.725	1.00		C
ATOM	648	C	VAL		87	119.146	60.342	15.513 15.893	1.00	32.90 32.72	0
ATOM	649	O CP	VAL VAL		87 87	119.515 117.637	61.458 58.952	16.950	1.00		C
ATOM	650 651	CB CG1	VAL		87	117.637	59.511	18.099	1.00		C
ATOM ATOM	652	CG2		A	87	116.196	58.796	17.394	1.00		C
ATOM	653	N	ARG	A	88	119.969	59.481	14.920	1.00	32.89	N
ATOM	654	CA	ARG		88	121.353	59.860	14.679	1.00		C
ATOM	655	C	ARG		88	121.385	61.104	13.813			C
ATOM	656	0			88	122.176	62.007	14.060	1.00		0
ATOM	657	CB	ARG	A	88	122.142	58.740	13.989	1.00	35.05	C
ATOM	658	CG	ARG	Α	88	122.412	57.533	14.861	1.00		C
ATOM	659	CD	ARG	A	88	123.375	56.540	14.195	1.00		C
ATOM	660	NE	ARG	A	88	123.796	55.505	15.141	1.00	43.00	N
ATOM	661	CZ	ARG	Α	88	123.038	54.493	15.550	1.00	43.55	С
ATOM	662	NH1	ARG	Α	88	123.515	53.610	16.421	1.00	43.61	N
ATOM	663	NH2	ARG	Α	88	121.819	54.340	15.060	1.00	45.10	N
MOTA	664	N	LYS		89	120.519	61.167	12.805	1.00	33.67	N
ATOM	665	CA	LYS	A	89	120.509	62.331	11.929	1.00	33.25	C
ATOM	666	C	LYS	Α	89	120.348	63.618	12.703	1.00	32.41	C
MOTA	667	0	LYS	Α	89	120.956	64.630	12.356	1.00	33.80	0

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ATOM	668	CB	LYS	Α	89	119.406	62.243	10.873	1.00 33.9	_	C
ATOM	669	CG	LYS	A	89	119.689	61.251	9.758	1.00 37.7		C
ATOM	670	CD	LYS	Α	89	118.720	61.454	8.596	1.00 39.4		C
MOTA	671	CE	LYS	A	89	118.999	60.487	7.453	1.00 39.8		C
ATOM	672	NZ	LYS	A	89	118.777	59.074	7.855	1.00 42.1		N
MOTA	673	N	VAL	A	90	119.539	63.597	13.755	1.00 30.9		N
ATOM	674	CA	VAL	A	90	119.346	64.811	14.526	1.00 29.4		C
ATOM	675	C	VAL	A	90	120.612	65.152	15.304	1.00 29.8		C
ATOM	676	0	VAL	Α	90	120.989	66.327	15.402	1.00 29.5		0
MOTA	677	CB	VAL		90	118.145	64.690	15.494	1.00 27.6		C
MOTA	678	CG1	VAL		90	118.020	65.958	16.335	1.00 26.3		C
ATOM	679	CG2	VAL	A	90	116.863	64.484	14.697	1.00 25.4		C
MOTA	680	N		A	91	121.280	64.132	15.839	1.00 30.2		N
ATOM	681	CA	LYS	A	91	122.502	64.372	16.607	1.00 30.6		C
ATOM	682	С		A	91	123.646	64.827	15.701	1.00 30.9		C
ATOM	683	0	LYS		91	124.487	65.625	16.105	1.00 32.0		0
ATOM	684	CB		A	91	122.921	63.113	17.371	1.00 28.8		C
ATOM	685	CG		A	91	121.862	62.546	18.307	1.00 28.2		C
ATOM	686	CD	LYS		91	121.288	63.582	19.281	1.00 28.8		C
MOTA	687	CE		A	91	122.332	64.181	20.217	1.00 27.5		C
ATOM	688	NZ			91	121.726	65.148	21.203	1.00 28.6		N
MOTA	689	N	ARG	A	92	123.664	64.329	14.470	1.00 32.0		И
ATOM	690	CA		A.	92	124.708	64.678	13.513	1.00 33.2		C
ATOM	691	C	ARG		92	124.491	66.031	12.856	1.00 33.9		C
ATOM	692	0	ARG		92	125.369	66.532	12.151	1.00 35.1		0
ATOM	693	CB	ARG		92	124.790	63.627	12.409	1.00 34.1 1.00 35.2		C
ATOM	694	CG		A	92	125.206	62.236	12.834 13.424	1.00 33.2		C
ATOM	695	CD	ARG		92	126.593	62.227 60.867	13.424	1.00 38.8		N
ATOM	696	NE		A A	92 92	127.118 128.235	60.524	14.142	1.00 40.3		C
ATOM	697 698	CZ	ARG		92	128.962	61.443	14.771	1.00 41.4		N
ATOM ATOM	699	NH2		A	92	128.622	59.254	14.148	1.00 43.1		N
ATOM	700	N	SER		93	123.325	66.625	13.075	1.00 34.1		N
ATOM	701	CA	SER		93	123.016	67.910	12.461	1.00 34.7		C
ATOM	702	C	SER		93	124.039	69.013	12.806	1.00 35.0		C
ATOM	703	0	SER		93	124.714	69.554	11.924	1.00 35.5		0
ATOM	704	СВ	SER		93	121.601	68.342	12.863	1.00 34.3		C
ATOM	705	OG	SER		93	121.171	69.453	12.097	1.00 33.8	34	0
ATOM	706	N	GLU		94	124.148	69.343	14.087	1.00 35.1	L6	N
ATOM	707	CA	GLU		94	125.077	70.367	14.545	1.00 35.9	95	C
ATOM	708	С	GLU	A	94	126.125	69.754	15.462	1.00 34.7	73	C
ATOM	709	0	GLU	Α	94	125.799	68.965	16.352	1.00 34.3	32	0
ATOM	710	СВ	GLU	Α	94	124.290	71.494	15.221	1.00 38.4	1	C
ATOM	711	CG	GLU		94	123.485	72.240	14.164	1.00 43.9	97	C
ATOM	712	CD	GLU		94	122.504	73.241	14.699	1.00 46.4	11	C
ATOM	713	OE1	GLU	A	94	122.889	74.140	15.485	1.00 49.2	20	0
ATOM	714	OE2	GLU	A	94	121.331	73.134	14.300	1.00 49.6	57	0

ATOM	715	N	ASN	A	95	127.387	70.118	15.239	1.00	33.02	N
ATOM	716	CA	ASN	Α	95	128.479	69.543	16.012	1.00	32.01	C
ATOM	717	C	ASN	Α	95	128.343	68.043	15.831	1.00	32.61	C
MOTA	718	0	ASN	Α	95	128.609	67.259	16.747	1.00	33.70	0
MOTA	719	CB	ASN	Α	95	128.372	69.913	17.492	1.00	30.02	C
ATOM	720	CG	ASN	Α	95	128.838	71.321	17.770	1.00	29.11	C
MOTA	721	OD1	ASN	A	95	128.954	71.729	18.923	1.00	29.21	0
ATOM	722	ND2	ASN	A	95	129.111	72.078	16.709	1.00	29.45	N
MOTA	723	N	GLY	Α	96	127.930	67.669	14.620	1.00	32.33	N
ATOM	724	CA	GLY	A	96	127.697	66.283	14.265	1.00	31.72	C
ATOM	725	C	GLY		96	128.798	65.309	14.592	1.00	31.80	С
ATOM	726	0	GLY		96	128.522	64.157	14.919	1.00		0
ATOM	727	N	VAL		97	130.044	65.759	14.494	1.00		N
ATOM	728	CA	VAL		97	131.176	64.893	14.779	1.00		C
ATOM	729	C	VAL		97	132.092	65.468	15.859	1.00		C
MOTA	730	0	VAL		97	133.265	65.111	15.944	1.00		0
ATOM	731	CB	VAL		97	131.988	64.591	13.483	1.00		C
ATOM	732	CG1	VAL		97	131.121	63.797	12.506	1.00		C
MOTA	733	CG2	VAL		97	132.467	65.888	12.836	1.00		C
ATOM	734	N	ILE		98	131.542	66.357	16.684	1.00		N
ATOM	735	CA	ILE		98	132.288	66.962	17.789		30.43	C
ATOM	736	C	ILE		98	132.855	65.806	18.637		29.83	C
ATOM	737	O CB	ILE		98	133.873	65.950	19.309	1.00		0
ATOM	738	CB CG1	ILE		98 98	131.352	67.877 68.633	18.652 19.712	1.00		C
ATOM ATOM	739 740	CG2	ILE		98	132.152 130.264	67.044	19.712	1.00		C
ATOM	741	CD1	ILE		98	133.036	69.704	19.171		27.65	C
ATOM	742	N	ILE	A	99	132.174	64.665	18.593	1.00	29.88	N
ATOM	743	CA	ILE		99	132.594	63.450	19.290	1.00		C
ATOM	744	C	ILE		99	132.126	62.306	18.397	1.00		C
ATOM	745	0	ILE		99	131.219	62.488	17.580		34.65	0
ATOM	746	СВ		A	99	131.937	63.285	20.675	1.00		C
ATOM	747	CG1	ILE	Α	99	130.431	63.074	20.522	1.00		С
ATOM	748	CG2	ILE	A	99	132.225	64.503	21.530	1.00	29.88	C
ATOM	749	CD1	ILE	A	99	129.736	62.684	21.811	1.00	29.26	C
ATOM	750	N	ASP	A	100	132.724	61.130	18.544	1.00	33.17	N
ATOM	751	CA	ASP	A	100	132.353	59.993	17.704	1.00	34.04	C
ATOM	752	C	ASP	A	100	132.387	60.391	16.234	1.00	33.78	C
ATOM	753	0	ASP	Α	100	131.398	60.248	15.520	1.00	34.06	0
MOTA	754	CB	ASP	A	100	130.956	59.485	18.050	1.00	35.98	C
ATOM	755	CG	ASP	A	100	130.844	59.045	19.490	1.00	39.21	C
ATOM	756	OD1	ASP	A	100	131.737	58.305	19.952	1.00	41.93	0
ATOM	757	OD2	ASP	A	100	129.861	59.425	20.159	1.00	41.57	0
MOTA	758	N	PRO			133.532	60.909	15.767		33.06	N
ATOM	759	CA	PRO			133.697	61.334	14.377		33.37	C
MOTA	760	C	PRO			133.763	60.179	13.382		32.86	C
MOTA	761	0	PRO	A	101	133.952	59.031	13.763	1.00	33.47	0

ATOM	762	CB	PRO A	101	135.007	62.112	14.438	1.00	33.67	С
ATOM	763	CG	PRO A	101	135.786	61.284	15.427	1.00	32.82	С
ATOM	764	CD	PRO A	101	134.765	61.193	16.526	1.00	32.08	С
ATOM	765	N	PHE A	102	133.589	60.488	12.104	1.00	33.11	N
ATOM	766	CA	PHE A	102	133.693	59.469	11.065	1.00	33.73	C
ATOM	767	C	PHE A	102	135.160	59.479	10.692	1.00	33.47	C
ATOM	768	0	PHE A	102	135.813	60.506	10.805	1.00	34.81	0
ATOM	769	CB	PHE A	102	132.876	59.832	9.817	1.00	34.53	C
ATOM	770	CG	PHE A	102	131.396	59.878	10.045	1.00	35.48	C
ATOM	771	CD1	PHE A	102	130.703	58.738	10.431	1.00	35.76	C
ATOM	772	CD2	PHE A	102	130.691	61.069	9.880	1.00	36.37	C
ATOM	773	CE1	PHE A	102	129.325	58.787	10.652	1.00	37.70	C
ATOM	774	CE2	PHE A	102	129.313	61.124	10.097	1.00	36.90	C
MOTA	775	CZ	PHE A	102	128.632	59.982	10.483	1.00	36.80	C
MOTA	776	N	PHE A	103	135.684	58.346	10.252	1.00	34.52	N
MOTA	777	CA	PHE A	103	137.081	58.283	9.858	1.00	35.62	C
ATOM	778	C	PHE A	103	137.394	57.001	9.104	1.00	36.66	C
ATOM	779	0		103	136.670	56.013	9.215	1.00	37.58	0
ATOM	780	CB		103	137.989	58.453	11.094	1.00	34.86	C
ATOM	781	CG		103	137.695	57.493	12.221		34.87	С
MOTA	782	CD1			138.125	56.168	12.167		34.75	C
ATOM	783			103	136.978	57.917	13.340		34.59	C
ATOM	784	CE1		103	137.846	55.281	13.213		34.04	C
ATOM	785	CE2		103	136.696	57.037	14.387		34.29	C
ATOM ATOM	786 787	CZ N	PHE A LEU A	103	137.130 138.453	55.720 57.036	14.322		33.77	C
ATOM	788	CA		104	138.872	55.879	8.307 7.532		37.81 39.19	N C
ATOM	789	C	LEU A		140.378	55.799	7.650	1.00		C
ATOM	790	o		104	141.005	56.725	8.159		40.69	0
ATOM	791	CB	LEU A		138.470	56.035	6.062		39.32	C
ATOM	792	CG		104	136.970	56.076	5.745		39.96	C
ATOM	793	CD1	LEU A		136.769	56.381	4.278	1.00		C
MOTA	794	CD2	LEU A	104	136.321	54.747	6.108		40.67	C
ATOM	795	N	THR A	105	140.957	54.697	7.184	1.00	41.82	N
ATOM	796	CA	THR A	105	142.402	54.502	7.249	1.00	42.36	C
ATOM	797	C	THR A	105	143.017	54.739	5.880	1.00	43.67	C
ATOM	798	0	THR A	105	142.305	54.885	4.892	1.00	42.24	0
ATOM	799	CB	THR A	105	142.764	53.062	7.696	1.00	42.10	C
ATOM	800	OG1	THR A	105	142.395	52.126	6.673	1.00	41.32	0
ATOM	801	CG2	THR A		142.038	52.712	8.990	1.00	40.95	C
ATOM	802	N		106	144.354	54.802	5.810	1.00		N
MOTA	803	CA	PRO A		145.017	55.018	4.521	1.00		С
ATOM	804	C	PRO A		144.730	53.792	3.643	1.00		C
MOTA	805	0	PRO A		144.611	53.881	2.419	1.00		0
ATOM	806	CB	PRO A		146.493	55.109	4.920	1.00		C
ATOM	807	CG	PRO A		146.422	55.617	6.356	1.00		C
ATOM	808	CD	PRO A	100	145.364	54.693	6.876	1.00	45.95	C

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MOTA	809	N	GLU	A	107	144.606	52.651	4.312	1.00	51.71	N
MOTA	810	CA	GLU	Α	107	144.347	51.375	3.670	1.00	52.95	C
MOTA	811	C	GLU	A	107	143.116	51.449	2.777	1.00	52.83	C
ATOM	812	0	GLU	Α	107	143.177	51.081	1.608	1.00	53.42	0
MOTA	813	CB	GLU	Α	107	144.138	50.302	4.739	1.00	55.06	С
MOTA	814	CG	GLU	Α	107	145.023	50.465	5.974	1.00	58.02	С
ATOM	815	CD	GLU	A	107	146.496	50.615	5.636	1.00	60.20	C
MOTA	816	OE1	GLU	A	107	146.863	51.634	5.009	1.00	61.69	0
MOTA	817	OE2	GLU	Α	107	147.288	49.713	5.995	1.00	62.09	0
ATOM	818	N	HIS	Α	108	142.003	51.919	3.339	1.00	52.97	N
ATOM	819	CA	HIS	Α	108	140.736	52.045	2.611	1.00	52.34	C
MOTA	820	C	HIS	A	108	140.913	52.647	1.232	1.00	52.01	C
MOTA	821	0	HIS	Α	108	141.923	53.284	0.942	1.00	52.35	0
ATOM	822	CB	HIS	Α	108	139.745	52.907	3.398	1.00	52.32	C
ATOM	823	CG	HIS	A	108	139.233	52.257	4.642	1.00	52.17	С
ATOM	824	ND1	HIS	Α	108	138.355	51.198	4.619	1.00	53.21	N
ATOM	825	CD2	HIS	A	108	139.495	52.498	5.947	1.00	52.86	C
MOTA	826	CE1	HIS	A	108	138.097	50.814	5.856	1.00	52.97	C
MOTA	827	NE2	HIS	A	108	138.778	51.587	6.681	1.00	52.99	N
ATOM	828	N	LYS	Α	109	139.915	52.443	0.387	1.00	51.80	N
ATOM	829	CA	LYS	Α	109	139.945	52.962	-0.967	1.00	52.18	C
ATOM	830	C	LYS	Α	109	139.137	54.260	-1.038	1.00	52.51	С
ATOM	831	0	LYS	A	109	138.083	54.380	-0.409	1.00	52.67	0
ATOM	832	CB	LYS	Α	109	139.370	51.922	-1.929	1.00	52.23	C
ATOM	833	N	VAL	A	110	139.637	55.233	-1.792	1.00	52.49	N
ATOM	834	CA	VAL	Α	110	138.942	56.509	-1.938	1.00		C
ATOM	835	С	VAL	A	110	137.447	56.292	-2.190	1.00	52.34	С
ATOM	836	0	VAL	A	110	136.619	57.135	-1.838	1.00	52.26	0
ATOM	837	CB	VAL	Α	110	139.572	57.355	-3.087	1.00		C
ATOM	838	CG1	VAL	Α	110	139.815	56.494	-4.295		52.21	С
ATOM	839	CG2	VAL	A	110	138.655	58.499	-3.467	1.00		C
ATOM	840	N	SER	Α	111	137.111	55.151	-2.785	1.00		N
ATOM	841	CA	SER	A	111	135.721	54.813	-3.079		52.79	С
ATOM	842	C	SER	Ą	111	134.908	54.753	-1.799		52.49	C
ATOM	843	0	SER	A	111	133.805	55.295	-1.729	1.00	52.98	0
ATOM	844	CB	SER	A	111	135.630	53.457	-3.785	1.00	52.72	C
ATOM	845	OG	SER	Α	111	136.290	53.485	-5.035		54.78	0
ATOM	846	N	GLU	Α	112	135.458	54.086	-0.789		51.81	N
ATOM	847	CA	GLU	A	112	134.776	53.944	0.488	1.00		C
ATOM	848	C	GLU	A	112	134.617	55.315	1.135	1.00		C
ATOM	849	0	GLU	A	112	133.658	55.563	1.861	1.00		0
ATOM	850	CB	GLU	Α	112	135.565	52.990	1.391	1.00		C
ATOM	851	CG			112	135.912	51.665	0.706		50.40	C
ATOM	852	CD			112	136.572	50.652	1.633		51.24	C
ATOM	853	OE1			112	135.884	50.113	2.524		51.42	0
ATOM	854	OE2			112	137.786	50.399	1.476		51.68	0
ATOM	855	N	ALA	A	113	135.555	56.213	0.845	1.00	52.48	N

#### **ALA A 113** 57.570 C ATOM ÇA 135.509 1.392 1.00 53.74 856 857 **ALA A 113** 134.348 58.346 0.769 1.00 53.99 C MOTA C 1.450 MOTA 858 ALA A 113 133.653 59.108 1.00 53.28 0 0 MOTA ALA A 113 136.834 58.294 1.129 C 859 1.00 53.95 CB 860 **GLU A 114** 134.148 N ATOM N 58.157 -0.5311.00 54.68 **GLU A 114** 58.824 1.00 55.97 C CA MOTA 861 133.055 -1.221862 **GLU A 114** -0.781 1.00 56.86 C **ATOM** C 131.743 58.169 **MOTA GLU A 114** 0 863 0 130.685 58.799 -0.811 1.00 56.32 **GLU A 114** -2.732 C MOTA 864 CB 133.228 58.705 1.00 55.49 **MOTA GLU A 115** N 865 N 131.813 56.904 -0.366 1.00 57.91 866 C **ATOM** CA **GLU A 115** 130.616 56.209 0.076 1.00 59.80 **ATOM** 867 **GLU A 115** 1.293 1.00 60.45 C C 130.052 56.916 **ATOM** 128.866 1.326 0 868 **GLU A 115** 57.239 1.00 60.90 0 C **ATOM** 869 **GLU A 115** 130.905 54.747 0.444 1.00 61.61 CB C 870 131.454 53.877 CG **GLU A 115** -0.683 1.00 63.71 ATOM 131.451 52.391 1.00 64.61 C ATOM 871 CD **GLU A 115** -0.333 1.00 64.98 872 OE1 GLU A 115 131.959 52.023 0.753 0 MOTA OE2 GLU A 115 873 MOTA 1.00 65.96 0 130.946 51.590 -1.152874 2.294 LEU A 116 130.898 57.161 1.00 61.14 N MOTA N C 875 LEU A 116 57.830 3.504 MOTA CA 1.00 62.15 130.433 C MOTA **LEU A 116** 130.306 59.343 3.372 1.00 62.60 876 C 877 **LEU A 116** 129.583 59.975 4.149 1.00 62.47 0 0 MOTA 4.719 C 878 CB LEU A 116 131.299 57.443 1.00 62.38 MOTA C 879 LEU A 116 132.827 MOTA CG 57.374 4.656 1.00 62.61 C 133.375 880 CD1 LEU A 116 58.712 4.232 1.00 64.03 MOTA 133.389 56.975 1.00 60.86 C CD2 LEU A 116 6.020 881 MOTA MSE A 117 130.988 2.388 1.00 63.07 N HETATM 882 N 59.929 **HETATM** MSE A 117 61.368 2.176 C 883 CA 130.877 1.00 63.64 C 129.512 61.672 884 C MSE A 117 1.578 1.00 62.94 **HETATM** 1.770 1.00 63.05 0 **HETATM** 885 0 MSE A 117 128.967 62.754 C 1.00 65.51 886 CB MSE A 117 131.948 61.892 1.218 HETATM 133.350 61.990 1.00 68.03 C MSE A 117 1.775 HETATM 887 CG SE 134.418 62.891 **HETATM** 0.609 1.00 71.39 888 SE MSE A 117 C 1.00 71.27 889 MSE A 117 135.980 62.908 1.489 **HETATM** CE N 60.715 0.837 1.00 62.33 890 N **GLN A 118** 128.967 MOTA Ç MOTA 1.00 62.54 891 CA **GLN A 118** 127.666 60.904 0.216 C 892 **GLN A 118** MOTA C 126.526 60.316 1.036 1.00 61.98 Q 893 **GLN A 118 ATOM** 125.375 60.725 0.884 1.00 61.93 $\mathsf{C}$ MOTA **GLN A 118** 127.664 60.341 -1.215 1.00 63.28 894 CB C 1.00 65.32 MOTA 895 **GLN A 118** CG 128.093 61.366 -2.289 C MOTA 896 CDGLN A 118 129.504 61.938 -2.088 1.00 66.65 О 897 ATOM OE1 GLN A 118 129.903 62.898 -2.7601.00 66.36 NE2 GLN A 118 -1.179N MOTA 898 130.266 61.340 1.00 67.13 ARG A 119 126.843 N MOTA 59.371 1.915 1.00 61.19 899 N C 125.819 **MOTA** 2.762 1.00 60.86 900 CA ARG A 119 58.770 C ATOM C 3.884 1.00 60.07 901 ARG A 119 125.467 59.751 4.159 0 1.00 60.05 902 0 ARG A 119 124.294 59.993 ATOM

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MOTA	903	CB	ARG	A	119	126.322	57.458	3.373	1.00 61.79	C
MOTA	904	CG	ARG	A	119	125.271	56.709	4.188	1.00 63.14	C
MOTA	905	CD	ARG	A	119	125.850	55.496	4.928	1.00 64.66	C
MOTA	906	NE	ARG	A	119	126.606	54.585	4.064	1.00 66.88	N
MOTA	907	CZ	ARG	A	119	126.125	53.988	2.973	1.00 67.98	C
MOTA	908	NH1	ARG	Α	119	124.870	54.195	2.584	1.00 68.24	N
ATOM	909	NH2	ARG	A	119	126.902	53.171	2.269	1.00 68.20	N
ATOM	910	N			120	126.491	60.318	4.520	1.00 58.88	N
ATOM	911	CA	TYR			126.284	61.257	5.618	1.00 57.61	C
ATOM	912	C			120	126.450	62.692	5.172	1.00 57.09	C
ATOM	913	0			120	126.409	63.612	5.989	1.00 57.12	0
ATOM	914	CB	TYR		120	127.257	60.974	6.766	1.00 57.42	C
ATOM	915	CG			120	127.192	59.552	7.269	1.00 57.82	C
ATOM	916	CD1			120	127.890	58.529	6.623	1.00 57.64	C
ATOM	917	CD2	TYR			126.365	59.214	8.337	1.00 57.74	C
ATOM	918	CEI			120	127.760	57.207	7.028	1.00 57.96	C
ATOM	919	CE2			120	126.226	57.895	8.748	1.00 58.11 1.00 58.31	C
ATOM	920	CZ OH			120 120	126.923 126.772	56.897 55.590	8.091 8.496	1.00 58.89	0
ATOM	921 922	N			121	126.638	62.875	3.872	1.00 56.83	N
ATOM ATOM	923	CA			121	126.818	64.200	3.293	1.00 56.77	C
ATOM	924	C	ARG			127.838	65.035	4.069	1.00 56.25	C
ATOM	925	0			121	127.479	65.797	4.966	1.00 56.28	o
ATOM	926	CB			121	125.486	64.927	3.233	1.00 57.20	C
MOTA	927	N			122	129.110	64.875	3.715	1.00 56.16	N
ATOM	928	CA			122	130.198	65.608	4.349	1.00 55.55	C
MOTA	929	C			122	131.305	65.938	3.339	1.00 55.54	C
ATOM	930	0			122	131.590	65.158	2.418	1.00 54.76	0
ATOM	931	СВ			122	130.793	64.811	5.536	1.00 55.74	С
ATOM	932	CG1	ILE			131.116	63.377	5.094	1.00 55.80	С
ATOM	933	CG2		A	122	129.830	64.836	6.717	1.00 54.89	C
ATOM	934	CD1	ILE	Α	122	131.738	62.510	6.186	1.00 55.41	C
ATOM	935	N	SER	A	123	131.915	67.106	3.535	1.00 54.93	N
ATOM	936	CA	SER	A	123	132.984	67.625	2.681	1.00 54.06	C
ATOM	937	C	SER	A	123	134.342	66.963	2.907	1.00 52.28	C
ATOM	938	0	SER	A	123	135.218	67.017	2.037	1.00 51.82	0
ATOM	939	CB	SER	A	123	133.121	69.139	2.902	1.00 55.93	C
ATOM	940	OG	SER	A	123	134.227	69.683	2.192	1.00 58.65	0
MOTA	941	N	GLY	A	124	134.516	66.353	4.077	1.00 50.22	И
MOTA	942	CA	GLY	A	124	135.780	65.706	4.383	1.00 47.42	C
ATOM	943	C	GLY	Α	124	135.668	64.500	5.297	1.00 45.07	C
ATOM	944	0	GLY	Α	124	134.709	64.357	6.055	1.00 44.24	0
ATOM	945	N			125	136.674	63.634	5.222	1.00 43.19	N
ATOM	946	CA			125	136.728	62.421	6.031	1.00 40.55	C
ATOM	947	C			125	138.068	62.319	6.761	1.00 38.76	C
ATOM	948	0			125	139.108	62.120	6.127	1.00 38.57	0
ATOM	949	CB	VAL	A	125	136.569	61.164	5.143	1.00 40.66	C

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ATOM	950	CG1	VAL	A	125	136.596	59.900	5.994	1.00	38.37	C
MOTA	951	CG2	VAL	Α	125	135.287	61.260	4.349	1.00	40.81	C
ATOM	952	N	PRO	Α	126	138.066	62.478	8.101	1.00	36.88	И
MOTA	953	CA	PRO	Α	126	139.315	62.379	8.853	1.00	35.19	C
ATOM	954	C	PRO	A	126	139.932	61.024	8.545	1.00	35.43	С
ATOM	955	0	PRO	Α	126	139.225	60.021	8.465	1.00	35.78	0
ATOM	956	CB	PRO	A	126	138.837	62.486	10.296	1.00	34.50	C
ATOM	957	CG	PRO	Α	126	137.674	63.417	10.171	1.00	34.23	C
ATOM	958	CD	PRO			136.951	62.747	9.026	1.00		C
ATOM	959	N			127	141.244	60.993	8.351		35.94	N
ATOM	960	CA	ILE		127	141.934	59.741	8.048	1.00		C
ATOM	961	C			127	142.914	59.398	9.158		34.68	C
ATOM	962	0	ILE	A	127	143.803	60.183	9.481	1.00		0
ATOM	963	CB			127	142.673	59.831	6.687	1.00		C
ATOM	964	CG1	ILE			141.653	60.068	5.574	1.00		C
ATOM	965	CG2	ILE		127	143.454	58.559	6.420	1.00		C
ATOM	966	CD1	ILE	A	127	140.587	59.008	5.506	1.00		C
ATOM	967	N	VAL			142.733	58.220	9.744	1.00		N
ATOM	968	CA	VAL			143.577	57.769	10.842	1.00		C
ATOM	969	C	VAL			144.430	56.582	10.430	1.00		0
ATOM	970	O	LAV		128 128	144.079 142.721	55.831 57.391	9.522 12.065	1.00		C
ATOM	971	CB CG1	VAL			142.721	58.590	12.486		33.76	C
ATOM ATOM	972 973	CG2	VAL			141.820	56.211	11.732		34.06	C
ATOM	974	N			129	145.556	56.416	11.732		37.38	N
MOTA	975	CA			129	146.467	55.332	10.798	1.00		C
ATOM	976	C			129	145.759	53.984	10.730		40.56	C
ATOM	977	0			129	145.765	53.196	9.967		41.74	0
ATOM	978	CB			129	147.666	55.398	11.743		42.72	C
ATOM	979	CG			129	148.817	54.478	11.383		46.98	C
ATOM	980	CD			129	150.030	54.715	12.260		48.82	C
ATOM	981	OE1			129	151.062	54.051	12.044		51.32	0
ATOM	982	OE2	GLU		129	149.952	55.568	13.165		50.14	0
ATOM	983	N	THR	A	130	145.138	53.733	12.057	1.00	41.09	N
ATOM	984	CA	THR	Α	130	144.414	52.487	12.309	1.00	41.55	С
ATOM	985	C	THR	A	130	143.047	52.806	12.908	1.00	42.57	C
ATOM	986	0	THR	A	130	142.936	53.676	13.769	1.00	42.54	0
ATOM	987	CB	THR	Α	130	145.185	51.596	13.311	1.00	41.31	C
MOTA	988	OG1	THR	A	130	146.399	51.125	12.716	1.00	42.29	0
ATOM	989	CG2	THR	A	130	144.359	50.423	13.721	1.00	42.50	C
ATOM	990	N	LEU	A	131	142.004	52.115	12.459	1.00	43.48	N
MOTA	991	CA	LEU	Α	131	140.673	52.358	13.010	1.00	44.46	C
MOTA	992	C	LEU	A	131	140.687	52.246	14.537	1.00	45.25	C
MOTA	993	0	LEU	A	131	139.931	52.937	15.223	1.00	45.86	0
MOTA	994	CB	LEU	A	131	139.653	51.354	12.463	1.00	43.91	C
MOTA	995	CG	LEU	A	131	139.245	51.387	10.993		44.06	C
MOTA	996	CD1	LEU	A	131	138.161	50.346	10.768	1.00	43.41	C

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ATOM	997	CD2	LEU	Α	131	138.724	52.767	10.616	1.00	43.19	С
ATOM	998	N	ALA	Α	132	141.547	51.370	15.058	1.00	45.26	N
MOTA	999	CA	ALA	Α	132	141.662	51.144	16.497	1.00	45.23	С
MOTA	1000	C	ALA	Α	132	142.550	52.169	17.192	1.00	45.48	C
MOTA	1001	0	ALA	Α	132	142.308	52.513	18.350	1.00	45.96	0
ATOM	1002	CB	ALA	Α	132	142.194	49.742	16.757	1.00	45.56	C
ATOM	1003	N	ASN	Α	133	143.576	52.647	16.490	1.00	45.46	N
MOTA	1004	CA	ASN	Α	133	144.501	53.631	17.046	1.00	45.65	C
ATOM	1005	С	ASN	Α	133	143.949	55.048	16.975	1.00	43.65	С
ATOM	1006	0	ASN	Α	133	144.187	55.858	17.869	1.00	42.55	0
ATOM	1007	CB	ASN	A	133	145.843	53.574	16.311	1.00	49.92	C
ATOM	1008	CG	ASN	Α	133	146.520	52.219	16.448	1.00	56.12	C
ATOM	1009	OD1	ASN	Α	133	146.006	51.203	15.966	1.00	58.65	0
MOTA	1010	ND2	ASN	Α	133	147.675	52.191	17.123	1.00	58.07	N
ATOM	1011	N	ARG	Α	134	143.216	55.339	15.907	1.00	40.92	N
ATOM	1012	CA	ARG	A	134	142.625	56.658	15.695	1.00	39.45	C
ATOM	1013	C	ARG	Α	134	143.613	57.821	15.694	1.00	38.37	C
ATOM	1014	0	ARG	Α	134	143.246	58.957	15.992	1.00	36.45	0
ATOM	1015	CB	ARG	Α	134	141.497	56.900	16.709	1.00	38.26	C
ATOM	1016	CG	ARG	Α	134	140.300	56.027	16.404	1.00	38.05	C
MOTA	1017	CD	ARG	Α	134	139.161	56.142	17.384	1.00	38.82	С
MOTA	1018	NE	ARG	Α	134	138.665	57.501	17.541	1.00	39.64	N
ATOM	1019	CZ	ARG	Α	134	137.458	57.789	18.023	1.00	41.06	C
MOTA	1020	NH1	ARG	Α	134	136.635	56.808	18.381	1.00	41.95	N
ATOM	1021	NH2	ARG	Α	134	137.079	59.052	18.177	1.00	40.31	N
MOTA	1022	N	LYS	Α	135	144.861	57.532	15.335	1.00	38.36	N
MOTA	1023	CA	LYS	Α	135	145.888	58.562	15.277	1.00	38.55	С
ATOM	1024	C	LYS	Α	135	145.654	59.286	13.962	1.00	38.13	C
ATOM	1025	0	LYS	Α	135	145.719	58.689	12.888	1.00	38.05	0
MOTA	1026	CB	LYS	Α	135	147.286	57.944	15.286	1.00	40.56	C
MOTA	1027	CG	LYS	A	135	148.357	58.915	15.766	1.00	44.66	C
MOTA	1028	CD	LYS	Α	135	149.768	58.434	15.482	1.00	46.81	C
MOTA	1029	CE	LYS	Α	135	150.058	58.514	13.995	1.00	48.72	C
MOTA	1030	NZ	LYS	A	135	151.483	58.202	13.692	1.00	51.64	N
ATOM	1031	N	LEU	Α	136	145.372	60.577	14.041	1.00	37.97	N
MOTA	1032	CA	LEU	A	136	145.083	61.351	12.842	1.00	37.65	С
MOTA	1033	C	LEU	Α	136	146.305	61.414	11.939	1.00	37.71	С
MOTA	1034	0	LEU	A	136	147.377	61.816	12.375	1.00	37.36	0
ATOM	1035	CB	LEU	Α	136	144.675	62.772	13.222	1.00	36.41	C
ATOM	1036	CG	LEU	A	136	143.745	63.524	12.278	1.00	36.38	C
ATOM	1037	CD1	LEU	A	136	143.847	64.993	12.619	1.00	37.47	C
ATOM	1038	CD2	LEU	Α	136	144.126	63.310	10.839	1.00	37.55	C
ATOM	1039	N	VAL	A	137	146.153	61.015	10.684	1.00	37.97	N
ATOM	1040	CA	VAL	A	137	147.270	61.082	9.755	1.00	38.94	C
ATOM	1041	C	VAL	A	137	146.934	62.066	8.654	1.00	39.72	C
ATOM	1042	0	VAL	A	137	147.820	62.555	7.953	1.00	41.11	0
ATOM	1043	CB	VAL	A	137	147.599	59.708	9.131	1.00	39.02	C

ATOM	1044	CG1	VAL	A	137	148.108	58.761	10.207	1.00	39.27	С
ATOM	1045	CG2	VAL	A	137	146.369	59.134	8.452	1.00	39.34	C
MOTA	1046	N	GLY	Α	138	145.647	62.366	8.510	1.00	39.89	N
ATOM	1047	CA	GLY	A	138	145.225	63.305	7.487	1.00	40.00	C
MOTA	1048	C	GLY		138	143.723	63.410	7.327	1.00	40.02	C
ATOM	1049	0	GLY			142.964	62.994	8.197		39.08	0
ATOM	1050	N			139	143.292	63.972	6.208	1.00	40.37	N
MOTA	1051	CA			139	141.873	64.123	5.951	1.00	42.42	C
MOTA	1052	C			139	141.619	64.260	4.462	1.00	44.67	C
ATOM	1053	0			139	142.292	65.033	3.784	1.00	47.01	0
ATOM	1054	CB			139	141.318	65.358	6.674	1.00		C
ATOM	1055	CG1	ILE		139	139.861	65.581	6.289	1.00	39.76	C
ATOM	1056	CG2	ILE		139	142.149	66.568	6.332	1.00	41.34	C
ATOM	1057	CD1			139	139.218	66.707	7.054		39.54	C
ATOM	1058	N			140	140.656	63.504	3.946	1.00	46.05	N
ATOM	1059	CA		A	140	140.339	63.579	2.524	1.00	46.92	C
ATOM	1060	C			140	139.150	64.519	2.348	1.00		C
ATOM	1061	0			140	138.198	64.483	3.133	1.00	46.73	0
MOTA	1062	CB			140	140.011	62.179	1.944		47.40	C
ATOM	1063	CG1	ILE	A	140	139.843	62.274	0.430	1.00	48.09	C
ATOM	1064	CG2	ILE		140	138.739	61.622	2.572	1.00	46.23	C
ATOM	1065	CD1		A		139.614	60.936	-0.222		49.16	C
ATOM	1066	N	THR			139.221	65.379	1.334		48.17	N
ATOM	1067	CA	THR			138.152	66.343	1.072		49.05	C
ATOM	1068	C	THR			137.655	66.341	-0.379		50.14	C
ATOM	1069	0	THR			138.275	65.739	-1.259	1.00		0
ATOM	1070	CB	THR			138.606	67.774	1.423		47.83	C
ATOM	1071	OG1	THR			139.757	68.119	0.640		46.02	0 C
ATOM	1072	CG2	THR			138.942	67.872	2.899		47.21	
ATOM	1073	N	ASN			136.528	67.021	-0.608		51.50	N C
ATOM	1074	CA	ASN			135.933	67.126	-1.939		52.02 52.18	C
ATOM	1075	C	ASN			137.001	67.465	-2.969			0
ATOM	1076	O	ASN			137.129	66.788	-3.996 -1.971		51.47 52.46	C
ATOM	1077	CB	ASN			134.855	68.212 67.903	-1.971 -1.070	1.00	53.06	C
ATOM	1078	CG	ASN			133.687 133.150	66.803	-1.070		53.57	0
ATOM	1079	OD1	ASN ASN			133.150	68.884	-0.279		53.81	N
ATOM	1080	ND2 N	ARG			137.756	68.525	-2.692	1.00		N
ATOM	1081					137.736	68.951	-3.584	1.00	54.02	C
ATOM	1082	CA C	ARG ARG			139.664	67.726	-3.944	1.00		C
ATOM	1083					140.061	67.543	-5.096	1.00		0
ATOM	1084	O	ARG					-2.902	1.00		C
ATOM	1085	CB N	ARG		143	139.695 139.915	70.014	-2.902		55.80	И
ATOM	1086					140.700	65.677	-3.152		56.78	C
ATOM	1087	CA C	ASP			139.969	64.713	-4.074		56.65	C
ATOM	1088				144	140.574	64.713	-4.959		56.82	0
ATOM	1089	O			144		64.119	-1.812		58.80	C
ATOM	1090	CB	HOP	A	144	140.969	ロセ・フブエ	-1.012	1.00	50.00	C

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ATOM	1091	CG	ASP	A	144	141.647	65.910	-0.810	1.00	61.29	C
MOTA	1092	OD1	ASP	A	144	141.895	65.471	0.337	1.00	61.86	0
MOTA	1093	OD2	ASP	Α	144	141.931	67.075	-1.171	1.00	63.17	0
HETATM	1094	N	MSE	Α	145	138.667	64.562	-3.870	1.00	56.83	N
HETATM	1095	CA	MSE	Α	145	137.880	63.639	-4.682		57.70	C
HETATM	1096	C			145	137.785	64.062	-6.151		57.60	C
HETATM	1097	0	MSE			137.748	63.206	-7.041		57.31	0
HETATM		CB			145	136.471	63.475	-4.085		59.38	C
HETATM		CG			145	136.437	62.951	-2.630		60.94	C
HETATM	1100	SE			145	137.129	61.269	-2.351		62.97	SE
HETATM	1101	CE			145	135.969	60.258	-3.296		61.65	C
ATOM	1102	N			146	137.750	65.371	-6.407		57.09	N C
ATOM	1103	CA			146	137.664 138.988	65.877 65.762	-7.782 -8.528		56.31 56.60	C
ATOM	1104	C O			146 146	139.012	65.623	-9.750		56.62	0
ATOM	1105 1106	СВ			146	137.156	67.326	-7.793		54.46	C
ATOM ATOM	1107	CG			146	137.130	67.413	-7.370		53.47	C
ATOM	1107	CD			146	135.700	68.816	-7.389		52.51	C
ATOM	1100	NE			146	135.796	69.724	-6.471		51.48	N
ATOM	1110	CZ			146	135.178	70.651	-5.747		50.40	C
ATOM	1111	NH1				135.878	71.435	-4.942		50.47	N
ATOM	1112	NH2			146	133.858	70.774	-5.804		49.06	N
ATOM	1113	N			147	140.087	65.813	-7.787		57.55	N
ATOM	1114	CA			147	141.414	65.682	-8.375	1.00	58.70	C
ATOM	1115	C			147	141.641	64.221	-8.736	1.00	59.46	C
ATOM	1116	0	PHE	Α	147	142.318	63.911	-9.718	1.00	58.95	0
ATOM	1117	CB	PHE	Α	147	142.485	66.148	-7.377	1.00	58.89	C
ATOM	1118	CG	PHE	A	147	143.894	65.843	-7.807	1.00	59.83	С
MOTA	1119	CD1	PHE	A	147	144.369	64.530	-7.822	1.00	60.00	C
ATOM	1120	CD2	PHE	A	147	144.741	66.865	-8.228	1.00	60.66	C
ATOM	1121	CE1	PHE	A	147	145.662	64.238	-8.252	1.00	60.08	С
MOTA	1122	CE2	PHE	A	147	146.040	66.587	-8.661	1.00	60.69	С
MOTA	1123	CZ	PHE	Α	147	146.500	65.268	-8.673	1.00	60.63	С
ATOM	1124	N	ILE	A	148	141.062	63.335	-7.928		60.40	N
ATOM	1125	CA			148	141.187	61.889	-8.103		62.00	C
ATOM	1126	С			148	140.398	61.335	-9.291		63.07	C
ATOM	1127	0			148	139.203	61.593	-9.441		62.54	0
ATOM	1128	CB			148	140.735	61.148	-6.823		61.84	C
ATOM	1129	CG1	ILE		148	141.583	61.602	-5.640		60.88	C
ATOM	1130	CG2			148	140.870	59.641	-7.006		61.40	C
ATOM	1131	CD1			148	141.125	61.022	-4.335		61.89	N
ATOM	1132	N Ca			149	141.083		-10.120 -11.299		65.13 67.10	C
ATOM	1133	CA			149	140.476 139.843		-11.299		68.06	C
ATOM	1134	С 0			149 149	139.843		-10.333		68.25	0
MOTA MOTA	1135 1136	CB			149	141.522		-12.399		67.81	C
	1137	OG			149	140.952		-13.539		70.51	0
ATOM	TT2 /	UG	SEK	A	エセジ	140.954	JJ.1/	13.339	1.00	, , , , , ,	•

ATOM	1138	N	ASP	A	150	140.	655	57.64	19	-10.552	1.	00	68.	94	N
ATOM	1139	CA	ASP	Α	150	140.	150	56.31	<b>L</b> 5	-10.231	1.	00	70.	14	С
ATOM	1140	С	ASP	Α	150	139.	976	56.06	55	-8.731	1.	00	70.	46	C
ATOM	1141	0	ASP	Α	150	140.	951	56.00	7	-7.976	1.	00	69.	92	0
ATOM	1142	CB	ASP	A	150	141.	060	55.24	ł 0	-10.832	1.	00	70.	63	С
ATOM	1143	CG	ASP	Α	150	140.	627	53.83	34	-10.454	1.	00	71.	26	С
ATOM	1144	OD1	ASP	Α	150	140.	813	53.44	17	-9.281	1.	00	71.	97	0
ATOM	1145	OD2	ASP	Α	150	140.	084	53.12	20	-11.323	1.	00	71.	75	0
ATOM	1146	N	TYR	Α	151	138.	720	55.90	)1	-8.320	1.	00	71.	24	N
ATOM	1147	CA	TYR	Α	151	138.	374	55.66	57	-6.924	1.	00	72.	22	С
ATOM	1148	С	TYR	A	151	138.	715	54.28	31	-6.383	1.	00	73.	14	С
ATOM	1149	0	TYR	Α	151	138.	725	54.07	73	-5.173	1.	00	73.	06	0
ATOM	1150	CB	TYR	Α	151	136.	886	55.95	3	-6.698	1.	00	71.	67	C
ATOM	1151	CG	TYR	Α	151	136.	524	57.42	22	-6.784	1.	00	72.	11	С
ATOM	1152	CD1	TYR	A	151	135.	227	57.86	50	-6.503	1.	00	71.	81	C
ATOM	1153	CD2	TYR	Α	151	137.	484	58.38	33	-7.117	1.	00	72.	04	C
MOTA	1154	CE1	TYR	Α	151	134.	896	59.21	18	-6.548	1.	00	71.	71	C
ATOM	1155	CE2	TYR	Α	151	137.	165	59.74	ł O	-7.165	1.	00	71.	99	C
MOTA	1156	CZ	TYR	A	151	135.	872	60.15	51	-6.880	1.	00	72.	05	С
ATOM	1157	OH	TYR	A	151	135.	558	61.49	92	-6.927	1.	00	72.	25	0
MOTA	1158	N	ASN	A	152	138.	995	53.33	34	-7.270	1.	00	74.	39	N
ATOM	1159	CA	ASN	Α	152	139.	336	51.98	32	-6.841	1.	00	75.	27	C
MOTA	1160	С	ASN	A	152	140.	814	51.93	37	-6.437	1.	00	75.	50	C
ATOM	1161	0	ASN	Α	152	141.	587	51.11	13	-6.932	1.	00	75.	85	0
MOTA	1162	СВ	ASN	Α	152	139.	064	50.98	37	-7.976	1.	00	75.	63	C
MOTA	1163	CG	ASN	Α	152	139.	219	49.53	39	-7.539	1.	00	76.	28	С
ATOM	1164	OD1	ASN	Α	152	139.	084	48.62	0 2	-8.350	1.	00	77.	14	0
MOTA	1165	ND2	ASN	Α	152	139.	497	49.32	29	-6.254	1.	00	76.	29	N
MOTA	1166	N	ALA	Α	153	141.	199	52.83	34	-5.536	1.	00	75.	52	N
MOTA	1167	CA	ALA	Α	153	142.	578	52.90	)6	-5.067	1.	00	75.	70	C
MOTA	1168	C	ALA	Α	153	142.	649	53.36	0	-3.609	1.	00	76.	00	C
MOTA	1169	0	ALA	Α	153 ·	141.	778	54.09	93	-3.133	1.	00	75.	58	0
ATOM	1170	CB	ALA	Α	153	143.	375	53.86	54	-5.951	1.	00	75.	45	C
ATOM	1171	N	PRO	A	154	143.	696	52.92	29	-2.880	1.	00	76.	38	N
MOTA	1172	CA	PRO	Α	154	143.	883	53.29	4	-1.474	1.	00	76.	60	C
MOTA	1173	C	PRO	A	154	143.	829	54.80	9	-1.297	1.	00	76.	99	C
MOTA	1174	0	PRO	A	154	144.	532	55.55	52	-1.984	1.	00	76.	89	0
ATOM	1175	CB	PRO	Α	154	145.	258	52.70	7	-1.164	1.	00	76.	34	C
MOTA	1176	CG	PRO	Α	154	145.	247	51.45	57	-2.012	1.	00	75.	98	C
ATOM	1177	CD	PRO	A	154	144.	813	52.07	76	-3.321	1.	00	76.	29	C
MOTA	1178	N	ILE	Α	155	142.	988	55.25	8	-0.372	1.	00	77.	55	N
MOTA	1179	CA	ILE	A	155	142.	818	56.68	31	-0.114	1.	00	78.	03	C
MOTA	1180	C	ILE	A	155	144.		57.34		0.208		00	78.		C
ATOM	1181	0	ILE	A	155	144.		58.55		0.337			78.		0
MOTA	1182	CB	ILE	A	155	141.		56.91		1.050			77.		C
ATOM	1183		ILE					58.39		1.151			78.		C
ATOM	1184	CG2	ILE	A	155	142.	450	56.43	33	2.349	1.	00	77.	56	С

ATOM	1185	CD1	ILE	A	155		140.407	5	8.690	2	2.160	1	.00	78.30	C
ATOM	1186	N	SER	A	156		145.190	5	6.523	(	0.330	1	00	79.20	N
ATOM	1187	CA	SER	A	156		146.534	5	7.002	(	0.631	1	00	80.33	С
ATOM	1188	C	SER	Α	156		147.172	5	7.678	- (	0.584	1	.00	81.04	С
ATOM	1189	0	SER	A	156		147.530	5	8.859	- (	0.535	1	.00	81.05	0
MOTA	1190	CB	SER	Α	156		147.408	5	5.828	:	1.079	1	.00	80.24	C
MOTA	1191	OG	SER	Α	156		148.735	5	6.249	:	1.337	1	00	80.63	0
MOTA	1192	N	GLU	Α	157		147.308	5	6.914	- :	1.667	1	.00	81.98	N
MOTA	1193	CA	GLU	Α	157		147.904	5	7.392	-2	2.913	1	.00	82.66	C
MOTA	1194	C	GLU	Α	157		147.458	5	8.803	- 3	3.271	1	.00	83.46	С
MOTA	1195	0	GLU	Α	157		148.228	5	9.588	- 3	3.821	1	00	84.07	0
MOTA	1196	CB	GLU	A	157		147.556	5	6.426	- 4	4.047	1	.00	82.38	C
MOTA	1197	CG	GLU	A	157		148.089	5	5.022	- 3	3.808	1	.00	82.83	C
MOTA	1198	CD	GLU	Α	157		147.657	5	4.028	- 4	4.865	1	.00	83.03	C
MOTA	1199	OE1	GLU	Α	157		148.095	5	2.857	- 4	4.790	1	.00	82.84	0
MOTA	1200	OE2	GLU	Α	157		146.877	5	4.415	- [	5.763	1	.00	83.32	0
ATOM	1201	N	HIS	Α	158		146.209	5	9.117	-2	2.952	1	.00	84.27	N
MOTA	1202	CA	HIS	Α	158		145.643	6	0.434	- 3	3.219	1	.00	84.85	C
MOTA	1203	С	HIS	A	158		145.351	6	0.997	-:	1.845	1	.00	84.86	C
MOTA	1204	0	HIS	A	158		144.307	6	0.702	- 3	1.263	1	.00	84.99	0
ATOM	1205	CB	HIS	A	158		144.344	6	0.285	- 4	4.004	1	.00	85.54	C
MOTA	1206	CG	HIS	A	158		144.473	5	9.414	- 5	5.212	1	.00	86.62	C
ATOM	1207	ND1	HIS	A	158		145.274	5	9.744	- 6	5.284	1	.00	87.28	N
ATOM	1208	CD2	HIS	Α	158	,	143.939	5	8.204	- 5	5.499	1	.00	87.21	C
MOTA	1209	CE1	HIS	A	158	,	145.227	5	8.773	- 7	7.180	1	.00	87.78	C
ATOM	1210	NE2	HIS	Α	158		144.424	5	7.828	- 6	5.727	1	.00	87.86	N
HETATM	1211	N	MSE	Α	159	,	146.256	6	1.812	- :	1.319	1	.00	84.93	N
HETATM	1212	CA	MSE	A	159		146.042	6	2.323	(	0.020	1	.00	85.49	C
HETATM	1213	C	MSE	Α	159		146.301	6	3.765	(	0.395	1	.00	84.25	C
HETATM	1214	0	MSE	A	159	,	146.445	6	4.646	- (	0.450	1	.00	84.36	0
HETATM	1215	CB	MSE	Α	159	,	146.789	6	1.430	(	0.998	1	.00	87.98	C
HETATM	1216	CG	MSE	A	159	,	145.862	6	0.499	=	1.708	1	.00	91.79	C
HETATM	1217	SE	MSE	Α	159	,	144.854	6	1.381	2	2.951	1	.00	96.88	SE
HETATM	1218	CE	MSE	A	159		143.842	6	0.037	3	3.500	1	.00	95.69	C
ATOM	1219	N	THR	Α	160		146.334	6	3.979	-	1.707	1	.00	82.42	N
MOTA	1220	CA	THR	A	160		146.554	6	5.285	2	2.294	1	.00	80.55	C
ATOM	1221	С	THR	Α	160	,	147.496	6	5.164	3	3.494	1	.00	79.27	C
ATOM	1222	0	THR	Α	160	,	147.094	6	5.408	4	4.636	1	.00	79.55	0
ATOM	1223	CB	THR	Α	160		145.220	6	5.902	2	2.763	1	.00	80.73	C
ATOM	1224	OG1	THR	Α	160		144.314	6	5.984	-	1.655	1	.00	80.34	0
ATOM	1225	CG2	THR	Α	160		145.444	6	7.296	3	3.327	1	.00	80.81	С
ATOM	1226.	N	SER	A	161		148.740	6	4.758	3	3.238	1	.00	77.07	N
ATOM	1227	CA	SER	A	161		149.739	6	4.647	4	4.304	1	.00	74.22	С
ATOM	1228	C	SER	A	161		150.444	6	5.995	4	1.396	1	.00	72.09	C
ATOM	1229	0	SER	A	161	,	151.661	6	6.101	4	4.203	1	.00	72.10	0
ATOM	1230	CB	SER	Α	161	,	150.761	6	3.542	4	4.005	1	.00	74.18	C
MOTA	1231	OG	SER	A	161		150.194	6	2.254	4	4.160	1	.00	73.46	0

ATOM	1232	N	GLU	A	162	149.649	67.023	4.678	1.00	68.49	N
ATOM	1233	CA	GLU	A	162	150.138	68.386	4.802	1.00	63.99	C
MOTA	1234	C	GLU	Α	162	150.457	68.572	6.275	1.00	59.31	С
ATOM	1235	0	GLU	Α	162	150.507	67.606	7.030	1.00	59.06	0
MOTA	1236	CB	GLU	Α	162	149.045	69.372	4.378	1.00	66.51	C
ATOM	1237	CG	GLU		162	148.309	68.983	3.088	1.00		С
ATOM	1238	CD	GLU			149.171	69.063	1.834	1.00		C
ATOM	1239	OE1	GLU			148.736	68.542	0.781	1.00		0
ATOM	1240	OE2	GLU			150.266	69.661	1.890	1.00		0
ATOM	1241	N	HIS	A		150.673	69.812	6.683	1.00		N
ATOM	1242	CA	HIS			150.977	70.095	8.075	1.00		C
ATOM	1243	C	HIS		163	149.629	70.167	8.775		45.82	C
ATOM	1244	O CB	HIS			148.922	71.167	8.680	1.00		0
ATOM	1245	CB CG	HIS HIS	A	163	151.724 152.253	71.430 71.721	8.187 9.555	1.00		C
ATOM ATOM	1246 1247	ND1			163	153.047	70.833	10.249	1.00		N
ATOM	1247	CD2	HIS		163	152.151	72.822	10.335	1.00		C
ATOM	1249	CE1	HIS	A	163	153.412	71.374	11.396	1.00		C
ATOM	1250	NE2	HIS	Α	163	152.883	72.582	11.472	1.00		N
ATOM	1251	N	LEU			149.271	69.096	9.467	1.00		N
ATOM	1252	CA	LEU			147.990	69.033	10.156	1.00		C
ATOM	1253	C	LEU			147.812	70.110	11.218		36.83	C
ATOM	1254	0			164	148.676	70.312	12.068	1.00		0
ATOM	1255	СВ	LEU	Α	164	147.800	67.641	10.781	1.00		С
MOTA	1256	CG	LEU	A	164	147.792	66.468	9.789	1.00	39.23	C
ATOM	1257	CD1	LEU	Α	164	147.637	65.153	10.537	1.00	39.06	С
MOTA	1258	CD2	LEU	A	164	146.665	66.650	8.788	1.00	38.45	C
ATOM	1259	N	VAL	A	165	146.689	70.817	11.139	1.00	33.58	N
ATOM	1260	CA	VAL	Α	165	146.356	71.858	12.103	1.00	30.38	C
ATOM	1261	C	VAL	A	165	145.268	71.266	12.996	1.00	29.08	C
MOTA	1262	0	VAL	A	165	144.223	70.841	12.514	1.00	28.47	0
ATOM	1263	CB	VAL			145.840	73.131	11.395	1.00	29.47	C
ATOM	1264	CG1	VAL			145.440	74.176	12.417	1.00		C
ATOM	1265	CG2	VAL			146.931	73.687	10.486	1.00	28.52	C
ATOM	1266	N	THR			145.515	71.228	14.301	1.00		N
ATOM	1267	CA	THR			144.551	70.641	15.219	1.00	27.43	C
ATOM	1268	C	THR			144.388	71.443	16.492	1.00		C
ATOM	1269	O	THR			145.090	72.427	16.723	1.00		0
ATOM	1270	CB	THR			144.978	69.212	15.652	1.00		C
ATOM	1271 1272	OG1 CG2	THR			146.169 145.251	69.289 68.333	16.447	1.00	27.74	0
ATOM ATOM	1272	N	ALA			143.251	70.991	14.437 17.321	1.00	29.11 26.41	N
		CA	ALA			143.451	71.612	18.608		25.08	C
ATOM ATOM	1274 1275	CA	ALA			143.104	70.494	19.630		24.35	C
ATOM	1275	0	ALA			142.641	69.368	19.282		23.64	0
ATOM	1277	CB	ALA			141.900	72.441	18.517		24.85	C
ATOM	1278	N	ALA			143.249	70.804	20.893		24.53	N
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ATOM	1279	CA	ALA	A	168	143.131	69.808	21.947	1.00	24.62	C
MOTA	1280	С	ALA	A	168	141.679	69.424	22.180	1.00	25.16	C
MOTA	1281	0	ALA	Α	168	140.770	70.186	21.873	1.00	25.69	0
MOTA	1282	CB	ALA			143.731	70.346	23.232	1.00		C
ATOM	1283	N	VAL			141.461	68.234	22.722	1.00		N
ATOM	1284	CA	VAL			140.111	67.784	23.018	1.00		C
ATOM	1285	C	VAL			139.504	68.792	23.993	1.00		C
ATOM	1286	0	VAL			140.173	69.253	24.917	1.00		0
ATOM	1287	CB	VAL		169	140.133	66.394	23.670	1.00		С
ATOM ATOM	1288 1289	CG1 CG2	VAL VAL		169 169	138.720 140.843	65.941 65.403	23.967 22.750	1.00		C
ATOM	1290	N	GLY			138.247	69.157	23.770	1.00		N
ATOM	1291	CA	GLY			137.594	70.107	24.652	1.00		C
ATOM	1292	C	GLY			137.638	71.578	24.268	1.00	30.17	C
ATOM	1293	0	GLY			137.060	72.400	24.972	1.00		0
ATOM	1294	N	THR			138.313	71.940	23.181	1.00	31.70	N
ATOM	1295	CA	THR			138.357	73.347	22.800	1.00		C
ATOM	1296	С	THR	A	171	136.942	73.680	22.353	1.00	32.68	C
ATOM	1297	0	THR	Α	171	136.318	72.891	21.636	1.00	33.39	0
ATOM	1298	CB	THR	Α	171	139.329	73.610	21.632	1.00	34.57	C
MOTA	1299	OG1	THR	A	171	138.728	73.204	20.402	1.00	38.21	0
MOTA	1300	CG2	THR	Α	171	140.606	72.818	21.823	1.00	34.01	C
ATOM	1301	N	ASP	A	172	136.432	74.831	22.780	1.00	32.59	N
ATOM	1302	CA	ASP	A	172	135.074	75.242	22.441	1.00	31.82	C
ATOM	1303	C	ASP			135.001	75.879	21.064	1.00	31.23	C
ATOM	1304	0			172	136.026	76.090	20.417			0
ATOM	1305	CB			172	134.553	76.209	23.497		33.64	C
ATOM	1306	CG	ASP		172	135.373	77.478	23.574	1.00	36.95	C
ATOM	1307				172	135.141	78.284	24.499		40.09	0
ATOM	1308	OD2 N			172	136.245	77.682	22.706		38.33	O N
ATOM ATOM	1309 1310	CA	LEU LEU		173	133.785 133.578	76.183 76.782	20.618 19.307	1.00	30.04 30.23	N C
ATOM	1311	C	LEU			134.226	78.152	19.168	1.00		C
ATOM	1312	0	LEU			134.694	78.521	18.094		29.98	0
ATOM	1313	СВ	LEU			132.084	76.893	19.009		29.27	C
ATOM	1314	CG			173	131.337	75.561	18.956	1.00	30.61	C
ATOM	1315	CD1	LEU			129.898	75.791	18.506			C
ATOM	1316	CD2	LEU	Α	173	132.036	74.623	17.988		31.30	C
MOTA	1317	N	GLU	A	174	134.264	78.898	20.262	1.00	32.92	N
ATOM	1318	CA	GLU	Α	174	134.840	80.230	20.249	1.00	35.64	C
ATOM	1319	C	GLU	A	174	136.335	80.202	19.925	1.00	36.13	C
ATOM	1320	0	GLU	Α	174	136.824	81.034	19.157	1.00	35.92	0
ATOM	1321	CB	GLU	A	174	134.617	80.896	21.604	1.00	38.56	C
ATOM	1322	CG	GLU	A	174	134.920	82.379	21.627	1.00	43.61	С
MOTA	1323	CD	GLU			134.812	82.960	23.025		46.61	C
ATOM	1324		GLU			133.771	82.735	23.687		48.37	0
ATOM	1325	OE2	GLU	A	174	135.765	83.644	23.457	1.00	48.22	0

ATOM	1326	N	THR	A	175	137.064	79.254	20.508	1.00	36.59	N
ATOM	1327	CA	THR	A	175	138.501	79.156	20.252	1.00	37.34	C
ATOM	1328	C	THR	A	175	138.805	78.488	18.913	1.00	35.61	C
MOTA	1329	0	THR	A	175	139.771	78.853	18.231	1.00	35.02	0
ATOM	1330	CB	THR	A	175	139.234	78.394	21.383	1.00	39.60	C
MOTA	1331	OG1	THR	A	175	140.618	78.248	21.031	1.00	42.14	0
MOTA	1332	CG2	THR	A	175	138.631	77.015	21.596	1.00	42.29	C
ATOM	1333	N	ALA	Α	176	137.969	77.519	18.537	1.00	34.40	N
ATOM	1334	CA	ALA	A	176	138.122	76.803	17.271	1.00	33.17	C
MOTA	1335	C	ALA	A	176	137.936	77.780	16.115	1.00	32.96	C
ATOM	1336	0	ALA			138.682	77.751	15.142	1.00		0
ATOM	1337	CB	ALA			137.106	75.680	17.182		31.85	C
ATOM	1338	N	GLU			136.937	78.649	16.234		33.83	N
ATOM	1339	CA	GLU			136.655	79.657	15.217	1.00		C
ATOM	1340	C	GLU			137.902	80.499	15.001	1.00		C
ATOM	1341	0	GLU			138.252	80.855	13.876		36.40	0
ATOM	1342	CB	GLU			135.526	80.570	15.679	1.00		C
ATOM	1343	CG	GLU			135.167	81.644	14.675	1.00		C
ATOM	1344	CD	GLU			134.092	82.584	15.185	1.00		C
ATOM	1345	OE1	GLU			133.594	83.395	14.378	1.00	41.80 42.06	0
ATOM	1346	OE2 N	GLU ARG			133.750	82.522 80.815	16.390 16.105			O N
ATOM ATOM	1347 1348	CA			178	138.563 139.779	81.605	16.103		35.48 35.76	C
ATOM	1349	CA			178	140.891	80.849	15.372	1.00		C
ATOM	1350	0			178	141.604	81.418	14.536	1.00		0
ATOM	1351	CB	ARG			140.180	81.888	17.541			C
ATOM	1352	CG	ARG			141.397	82.765	17.737		41.57	C
ATOM	1353	CD			178	141.581	82.930	19.229	1.00		C
ATOM	1354	NE	ARG			140.343	83.441	19.807	1.00		N
ATOM	1355	CZ	ARG	Α	178	139.911	83.163	21.031	1.00	48.54	C
ATOM	1356	NH1	ARG	Α	178	140.618	82.369	21.828	1.00	48.33	N
ATOM	1357	NH2	ARG	Α	178	138.759	83.675	21.454	1.00	49.63	N
MOTA	1358	N	ILE	Α	179	141.034	79.564	15.680	1.00	32.20	N
MOTA	1359	CA	ILE	A	179	142.070	78.763	15.046	1.00	30.42	C
ATOM	1360	C	ILE	Α	179	141.753	78.583	13.556	1.00	30.40	C
ATOM	1361	0	ILE	Α	179	142.626	78.716	12.707	1.00	30.21	0
ATOM	1362	CB	ILE	Α	179	142.201	77.380	15.744	1.00	29.67	C
ATOM	1363	CG1	ILE	Α	179	142.477	77.585	17.239	1.00	28.14	C
ATOM	1364	CG2	ILE	A	179	143.327	76.560	15.100	1.00	27.22	C
ATOM	1365	CD1			179	142.463	76.313	18.067	1.00	26.32	C
ATOM	1366	N			180	140.499	78.292	13.240	1.00		N
ATOM	1367	CA			180	140.102	78.107	11.849	1.00		C
ATOM	1368	C	LEU			140.345	79.366	11.025		31.52	C
ATOM	1369	0	LEU			140.713	79.290	9.856		30.22	0
ATOM	1370	CB	LEU			138.623	77.713	11.769		31.47	C
ATOM	1371	CG	LEU			138.242	76.307	12.247		30.67	C
ATOM	1372	CDI	LEU	A	180	136.735	76.209	12.402	1.00	32.24	С

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MOTA	1373	CD2	LEU	A	180	138.752	75.273	11.256	1.00	30.35	C
ATOM	1374	N	HIS	A	181	140.139	80.524	11.642	1.00	33.19	N
MOTA	1375	CA	HIS	A	181	140.340	81.790	10.953	1.00	35.20	С
MOTA	1376	C	HIS	A	181	141.806	82.126	10.694	1.00	36.88	С
MOTA	1377	0	HIS	Α	181	142.149	82.651	9.631	1.00	36.71	0
ATOM	1378	CB	HIS	A	181	139.665	82.930	11.728	1.00	35.30	С
MOTA	1379	CG			181	138.169	82.952	11.590	1.00	36.21	C
MOTA	1380	ND1	HIS			137.379	83.913	12.186	1.00	35.97	N
MOTA	1381	CD2	HIS		181	137.323	82.143	10.906	1.00	35.37	C
MOTA	1382	CE1	HIS			136.113	83.695	11.875	1.00	35.54	С
ATOM	1383	NE2	HIS			136.052	82.628	11.100	1.00		N
ATOM	1384	N			182	142.677	81.827	11.650	1.00	38.45	N
ATOM	1385	CA	GLU			144.092	82.123	11.476	1.00	39.85	C
MOTA	1386	C			182	144.760	81.234	10.436	1.00	38.65	C
MOTA	1387	0			182	145.780	81.608	9.861	1.00		0
MOTA	1388	CB			182	144.849	81.960	12.790		43.25	C
MOTA	1389	CG	GLU			146.348	82.210	12.626	1.00		C
ATOM	1390	CD			182	147.159	81.826	13.855	1.00	53.96	C
MOTA	1391	OE1	GLU		182	148.394	82.037	13.826			0
MOTA	1392	OE2	GLU			146.570	81.310	14.840	1.00	55.87	0
MOTA	1393	N	HIS			144.183	80.062	10.196	1.00	37.07	N
MOTA	1394	CA			183	144.758	79.101	9.262			C
MOTA	1395	C			183	143.963	78.982	7.973		36.75	C
ATOM	1396	0			183	144.223	78.101	7.146		36.54	0
ATOM	1397	CB			183	144.846	77.743	9.955		34.77	C
ATOM	1398	CG			183	145.625	77.778	11.235	1.00		C
ATOM	1399		HIS			147.004	77.752	11.271		35.14	N
ATOM	1400		HIS			145.220	77.922	12.519		33.17	C
ATOM	1401		HIS			147.412	77.878	12.521		34.52	C
ATOM	1402	NE2	HIS			146.349	77.985	13.297		34.67	N
ATOM	1403	N			184	142.997	79.880	7.807		37.38	N
ATOM	1404	CA			184	142.147	79.887	6.622		37.97	C
ATOM	1405	C			184	141.668	78.485	6.287		36.24	C
ATOM	1406	0			184	141.815	78.034	5.157		34.85	0
ATOM	1407	CB			184	142.897	80.473	5.420		40.41	C C
ATOM	1408	CG			184	143.263	81.954	5.537		44.51	
ATOM	1409	CD			184	144.039	82.396	4.299		49.58	C
ATOM	1410	NE			184	143.298	82.080	3.076		54.75	N
ATOM	1411	CZ			184	143.764	82.247	1.839		57.24	C
ATOM	1412		ARG			144.987	82.733	1.640		58.91	N
ATOM	1413		ARG			143.007	81.923	0.796		58.17	N
ATOM	1414	N CA			185	141.107	77.797	7.279		34.80	N
ATOM	1415	CA			185	140.589	76.447	7.079		34.18	C
ATOM	1416	C			185	139.142	76.323	7.542		34.03	C
ATOM	1417	O			185	138.648	77.157	8.299		32.73	0
ATOM	1418	CB			185	141.433	75.387	7.822		33.30	C
ATOM	1419	CG1	ILE	A	<b>T Q D</b>	141.606	75.789	9.286	T.00	32.78	C

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MOTA	1420	CG2	ILE	A	185	142.767	75.198	7.114	1.00	33.84	С
MOTA	1421	CD1	ILE	Α	185	142.348	74.769	10.120	1.00	32.98	C
MOTA	1422	N	GLU	Α	186	138.467	75.273	7.086	1.00	34.75	N
MOTA	1423	CA	GLU	A	186	137.075	75.057	7.450	1.00	35.67	C
ATOM	1424	C	GLU	A	186	136.815	73.918	8.430	1.00	34.22	C
MOTA	1425	0	GLU	Α	186	135.795	73.926	9.110	1.00	34.10	0
MOTA	1426	CB	GLU	Α	186	136.242	74.863	6.187	1.00	38.26	C
MOTA	1427	CG	GLU	A	186	136.193	76.098	5.321	1.00		С
MOTA	1428	CD	GLU	A	186	135.354	75.899	4.080	1.00		С
MOTA	1429	OE1	GLU	A	186	135.168	76.884	3.328	1.00		0
MOTA	1430	OE2	GLU	A	186	134.880	74.761	3.848	1.00		0
MOTA	1431	N	LYS			137.721	72.942	8.496	1.00		N
MOTA	1432	CA	LYS			137.590	71.805	9.422	1.00		C
MOTA	1433	C				138.755	71.785	10.411	1.00		C
MOTA	1434	0			187	139.917	71.876	10.012	1.00		0
MOTA	1435	CB				137.565	70.456	8.674	1.00		C
ATOM	1436	CG		A		136.246	70.067	8.021	1.00		C
ATOM	1437	CD	LYS		187	135.898	70.942	6.827	1.00		C
ATOM	1438	CE	LYS			136.826	70.688	5.634	1.00		C
ATOM	1439	NZ	LYS			136.695	69.309	5.064	1.00		N
ATOM	1440	N	LEU			138.451	71.654	11.697	1.00		N
ATOM	1441	CA			188	139.500	71.619	12.714		26.39	C
ATOM	1442	C			188	139.439	70.340	13.533	1.00		C
ATOM	1443	O			188	138.629	70.228	14.448		26.83	O C
ATOM	1444	CB			188	139.364	72.812	13.658 14.804		25.30 25.12	C
ATOM	1445	CG CD1			188 188	140.373 141.779	72.899 73.097	14.245		24.40	C
ATOM ATOM	1446 1447	CD2			188	140.001	74.055	15.709		25.16	C
ATOM	1448	N			189	140.001	69.360	13.703		25.63	N
ATOM	1449	CA			189	140.327	68.088	13.949		25.71	C
ATOM	1450	C			189	140.736	68.292	15.405		25.74	C
ATOM	1451	0			189	141.598	69.120	15.704		25.84	0
ATOM	1452	СВ			189	141.372	67.284	13.179		25.33	C
ATOM	1453	CG			189	141.290	67.884	11.789		26.79	C
ATOM	1454	CD			189	141.316	69.339	12.163		25.56	C
ATOM	1455	N			190	140.104	67.546	16.306		26.25	N
ATOM	1456	CA			190	140.414	67.623	17.729	1.00	25.59	С
MOTA	1457	С	LEU	Α	190	141.105	66.314	18.076	1.00	26.60	С
MOTA	1458	0	LEU	A	190	140.565	65.227	17.833	1.00	26.44	0
ATOM	1459	CB	LEU	A	190	139.139	67.777	18.557	1.00	24.57	C
MOTA	1460	CG	LEU	Α	190	138.236	68.966	18.223	1.00	24.56	C
MOTA	1461	CD1	LEU	A	190	137.073	68.992	19.206	1.00	24.55	C
MOTA	1462	CD2	LEU	A	190	139.026	70.266	18.309	1.00	24.12	С
MOTA	1463	N	VAL	A	191	142.303	66.411	18.639	1.00	27.41	N
MOTA	1464	CA	VAL	A	191	143.069	65.223	18.984	1.00	26.97	C
MOTA	1465	C	VAL	A	191	143.589	65.339	20.403	1.00	27.90	C
MOTA	1466	0	VAL	A	191	143.693	66.442	20.929	1.00	27.60	0

#### VAL A 191 65.065 **ATOM** 1467 CB 144.267 18.024 1.00 26.42 C C CG1 VAL A 191 143.769 65.012 1.00 25.29 ATOM 16.585 1468 CG2 VAL A 191 1.00 25.51 C **ATOM** 145.236 66.225 18.196 1469 1.00 29.14 **ASP A 192** 143.895 64.211 21.039 N ATOM 1470 N 1.00 29.84 MOTA 1471 ASP A 192 64.285 22.381 C 144.449 CA 22.287 1.00 29.63 C **ASP A 192** 145.958 ATOM 1472 64.466 C **ASP A 192** 146.528 64.519 21.190 1.00 28.08 0 ATOM 1473 1.00 29.60 **ASP A 192** 144.117 63.049 23.237 C MOTA 1474 CB ASP A 192 144.600 61.748 22.633 1.00 29.23 C MOTA 1475 CG OD1 ASP A 192 145.698 1.00 29.02 61.693 22.045 0 ATOM 1476 OD2 ASP A 192 1.00 32.38 0 143.880 60.751 22.791 MOTA 1477 **ASN A 193** 146.595 64.564 23.447 1.00 30.91 **ATOM** 1478 N N **ASN A 193** 64.774 1.00 32.71 C **ATOM** 1479 23.525 CA 148.033 **ASN A 193** 1.00 33.29 C ATOM 1480 148.867 63.703 22.840 C **ASN A 193** 150.085 22.720 1.00 33.19 0 63.832 1481 0 MOTA 148.440 64.909 MOTA **ASN A 193** 24.987 1.00 34.40 C 1482 CB C **ASN A 193** 149.093 66.230 25.272 1.00 35.62 **ATOM** 1483 CG 24.901 MOTA OD1 ASN A 193 148.574 67.279 1.00 37.39 0 1484 ND2 ASN A 193 66.193 25.938 1.00 38.24 1485 150.236 N MOTA MOTA 1486 SER A 194 148.203 62.652 22.376 1.00 33.93 N N 1.00 33.58 C 1487 SER A 194 148.879 MOTA CA 61.564 21.691 C MOTA SER A 194 148.552 61.538 20.209 1.00 32.89 1488 C MOTA 1489 SER A 194 1.00 33.82 0 148.921 60.592 19.522 0 **ATOM** SER A 194 C 1490 CB 148.494 60.227 22.323 1.00 34.55 SER A 194 0 149.014 60.139 23.643 1.00 37.23 MOTA 1491 OG MOTA 1492 **GLY A 195** 1.00 31.61 N N 147.861 62.566 19.717 **GLY A 195** 1.00 29.53 1493 62,613 C 147.505 18.305 MOTA CA 1.00 29.01 MOTA 1494 **GLY A 195** 17.930 C C 146.314 61.738 **ATOM GLY A 195** 61.483 16.756 0 1495 0 146.060 1.00 28.69 ARG A 196 145.578 61.274 18.932 1.00 28.50 N **ATOM** 1496 N CA C MOTA 1497 ARG A 196 144.412 60.441 18.696 1.00 28.40 1498 ARG A 196 143.202 61.325 C 18.409 1.00 27.84 MOTA C MOTA 142.906 1.00 27.44 0 1499 0 ARG A 196 62.253 19.161 59.560 19.922 C 1500 ARG A 196 ATOM 144.152 1.00 30.26 CB C ARG A 196 19.887 1.00 31.17 MOTA 1501 CG 142.860 58.752 C MOTA 1502 1.00 33.36 ARG A 196 142.663 58.063 21.216 CD MOTA 1503 NE ARG A 196 141.253 57.834 21.501 1.00 38.05 N 57.008 140.471 C 20.822 ARG A 196 1.00 39.60 MOTA 1504 CZNH1 ARG A 196 140.966 1505 56.321 19.808 1.00 42.65 N MOTA NH2 ARG A 196 139.192 21.152 N MOTA 1506 56.877 1.00 40.11 MOTA 1507 LEU A 197 142.510 61.030 17.313 1.00 27.30 N N 16.904 C **ATOM** 1.00 27.90 1508 LEU A 197 141.332 61.792 CA C LEU A 197 140.125 61.586 17.825 1.00 27.27 MOTA 1509 C 0 MOTA 1510 1.00 26.44 0 LEU A 197 139.661 60.460 17.997 C LEU A 197 61.394 15.488 1.00 26.93 MOTA 1511 140.928 CB C LEU A 197 MOTA 1512 1.00 26.60 139.711 62.116 14.912 CG C **ATOM** 1513 CD1 LEU A 197 140.067 63.569 14.604 1.00 24.69

ATOM	1514	CD2	LEU	A	197	139.268	61.396	13.637	1.00 26.73	C
ATOM	1515	N	SER	Α	198	139.610	62.674	18.393	1.00 26.75	N
ATOM	1516	CA	SER	A	198	138.452	62.585	19.277	1.00 27.12	C
ATOM	1517	С	SER	A	198	137.202	63.163	18.635	1.00 27.80	C
ATOM	1518	0	SER	Α	198	136.093	62.836	19.038	1.00 29.05	0
ATOM	1519	CB	SER	Α	198	138.703	63.331	20.581	1.00 27.75	C
ATOM	1520	OG	SER	Α	198	138.836	64.720	20.352	1.00 28.75	0
ATOM	1521	N	GLY	Α	199	137.380	64.035	17.647	1.00 27.55	N
ATOM	1522	CA	GLY	Α	199	136.240	64.641	16.990	1.00 26.57	C
ATOM	1523	С	GLY	Α	199	136.681	65.739	16.050	1.00 26.31	C
ATOM	1524	0	GLY	Α	199	137.874	65.905	15.814	1.00 26.69	0
ATOM	1525	N	LEU	Α	200	135.727	66.502	15.534	1.00 25.94	N
ATOM	1526	CA	LEU	Α	200	136.031	67.564	14.590	1.00 26.36	С
ATOM	1527	С	LEU	Α	200	135.040	68.719	14.635	1.00 25.73	C
ATOM	1528	0	LEU			133.839	68.505	14.730	1.00 24.48	0
ATOM	1529	СВ	LEU			136.102	66.966	13.173	1.00 28.24	C
ATOM	1530	CG	LEU	A	200	136.355	67.848	11.938	1.00 30.21	С
ATOM	1531	CD1		A	200	136.938	67.013	10.809	1.00 30.99	C
ATOM	1532	CD2	LEU	A	200	135.063	68.517	11.506	1.00 32.05	С
ATOM	1533	N	ILE		201	135.567	69.942	14.584	1.00 26.65	N
ATOM	1534	CA			201	134.759	71.160	14.591	1.00 27.46	C
ATOM	1535	C	ILE			134.862	71.811	13.205	1.00 29.38	C
ATOM	1536	0			201	135.944	71.871	12.604	1.00 29.46	0
ATOM	1537	CB			201	135.244	72.150	15.660	1.00 26.43	C
ATOM	1538	CG1			201	135.132	71.508	17.047	1.00 27.07	C
ATOM	1539	CG2			201	134.417	73.423	15.595	1.00 27.71	C
ATOM	1540	CD1			201	135.613	72.380	18.218	1.00 24.40	C
ATOM	1541	N	THR			133.728	72.296	12.703	1.00 30.75	N
ATOM	1542	CA	THR			133.660	72.906	11.379	1.00 32.31	C
ATOM	1543	C	THR			133.207	74.358	11.414	1.00 32.21	C
ATOM	1544	0	THR			132.650	74.825	12.405	1.00 31.14	0
ATOM	1545	СВ	THR			132.691	72.113	10.472	1.00 33.40	C
ATOM	1546	OG1	THR			133.146	70.760	10.361	1.00 36.21	0
ATOM	1547	CG2			202	132.632	72.718	9.080	1.00 36.76	C
	1548	N			203	133.452	75.066	10.315	1.00 33.57	N
ATOM		CA			203	133.452	76.461	10.196	1.00 33.87	C
ATOM	1549				203	131.531	76.544	10.244	1.00 33.07	C
ATOM	1550	C				130.978	77.430	10.887	1.00 34.10	0
ATOM	1551	O			203		77.430	8.877	1.00 33.87	C
ATOM	1552	CB			203	133.621		8.869	1.00 33.87	C
ATOM	1553	CG1			203	133.384	78.589	7.664	1.00 33.89	C
ATOM	1554	CG2			203	132.995	76.434			C
ATOM	1555	CD1			203	134.165	79.321	9.950	1.00 34.38	N
ATOM	1556	N			204	130.852	75.605	9.587	1.00 34.73	C
ATOM	1557	CA			204	129.388	75.578	9.588	1.00 36.17	C
ATOM	1558	C			204	128.838	75.505	11.014	1.00 36.21	0
ATOM	1559	0			204	127.916	76.243	11.360	1.00 36.39	C
ATOM	1560	CB	LYS	Α	204	128.857	74.386	8.782	1.00 38.20	C

ATOM	1561	CG	LYS	Α	204	129.058	74.473	7.284	1.00	41.84	C
ATOM	1562	CD	LYS	Α	204	130.529	74.521	6.918	1.00	47.07	C
ATOM	1563	CE	LYS	Α	204	130.723	74.686	5.410	1.00	50.14	C
ATOM	1564	NZ	LYS	Α	204	132.165	74.843	5.037	1.00	52.54	N
ATOM	1565	N	ASP	Α	205	129.391	74.611	11.834	1.00	36.04	N
MOTA	1566	CA	ASP	Α	205	128.950	74.472	13.224	1.00	36.48	C
MOTA	1567	С	ASP	Α	205	128.922	75.830	13.886	1.00	35.58	C
MOTA	1568	0	ASP	Α	205	127.995	76.162	14.624	1.00	36.04	0
ATOM	1569	CB	ASP		205	129.909	73.606	14.036	1.00	38.56	C
MOTA	1570	CG	ASP			129.957	72.189	13.569	1.00	40.22	С
ATOM	1571	OD1	ASP	Α	205	130.764	71.431	14.141	1.00	41.68	0
ATOM	1572	OD2	ASP	Α	205	129.196	71.833	12.644	1.00	42.07	0
MOTA	1573	N	ILE	Α	206	129.970	76.604	13.637	1.00	34.72	N
MOTA	1574	CA	ILE	Α	206	130.092	77.933	14.209	1.00	34.65	C
MOTA	1575	С	ILE	Α	206	129.025	78.854	13.628	1.00	34.32	C
ATOM	1576	0	ILE	Α	206	128.466	79.693	14.326	1.00	34.37	0
ATOM	1577	CB	ILE	Α	206	131.494	78.485	13.933	1.00	34.36	С
ATOM	1578	CG1	ILE	Α	206	132.522	77.495	14.486	1.00	35.28	C
MOTA	1579	CG2	ILE	Α	206	131.665	79.849	14.575	1.00	34.33	C
MOTA	1580	CD1	ILE	A	206	133.948	77.829	14.164	1.00	36.11	C
MOTA	1581	N	GLU	A	207	128.732	78.669	12.348	1.00	34.71	N
ATOM	1582	CA	GLU	A	207	127.737	79.479	11.671	1.00	35.12	C
ATOM	1583	C	GLU	A	207	126.326	79.117	12.120	1.00	34.48	C
MOTA	1584	0	GLU	Α	207	125.462	79.991	12.222	1.00	34.41	0
MOTA	1585	CB	GLU	A	207	127.864	79.302	10.160	1.00	36.78	C
ATOM	1586	CG	GLU	A	207	129.266	79.555	9.649	1.00	39.03	C
ATOM	1587	CD	GLU	A	207	129.353	79.532	8.139	1.00	41.21	С
ATOM	1588	OE1	GLU	A	207	128.871	78.550	7.519	1.00	41.65	0
ATOM	1589	OE2	GLU	Α	207	129.918	80.497	7.575	1.00	43.02	0
ATOM	1590	N	LYS	Α	208	126.090	77.835	12.392	1.00	32.85	N
MOTA	1591	CA	LYS	Α	208	124.770	77.396	12.838	1.00	31.74	C
MOTA	1592	C	LYS	Α	208	124.462	77.924	14.223	1.00	30.98	C
MOTA	1593	0	LYS	A	208	123.306	78.107	14.577	1.00	31.35	0
MOTA	1594	CB	LYS	A	208	124.668	75.874	12.824	1.00	31.08	C
MOTA	1595	CG	LYS	Α	208	124.654	75.281	11.429	1.00	32.19	C
ATOM	1596	CD	LYS	A	208	124.629	73.769	11.485	1.00	32.69	C
ATOM	1597	CE	LYS	Α	208	124.667	73.162	10.102	1.00	33.70	С
ATOM	1598	NZ	LYS	A	208	124.676	71.678	10.190	1.00	35.26	N
ATOM	1599	N	VAL	Α	209	125.499	78.172	15.010	1.00	31.31	N
MOTA	1600	CA	VAL	Α	209	125.305	78.713	16.346	1.00	31.66	C
ATOM	1601	C	VAL	A	209	124.770	80.136	16.169	1.00	32.91	C
ATOM	1602	0	VAL			123.948	80.614	16.953		33.61	0
ATOM	1603	CB	VAL	A	209	126.634	78.733	17.137	1.00	30.78	C
ATOM	1604	CG1	VAL	A	209	126.430	79.378	18.494		30.86	C
ATOM	1605	CG2	VAL			127.151	77.323	17.307		29.89	С
ATOM	1606	N	ILE			125.236	80.807	15.120		33.36	N
ATOM	1607	CA	ILE	A	210	124.791	82.164	14.822	1.00	34.24	C

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ATOM 1608 C ILE A 210 123.57 82.142 14.291 1.00 33.53 ATOM 1609 O ILE A 210 125.810 82.843 14.798 1.00 32.71 ATOM 1610 CB ILE A 210 127.909 83.101 14.378 1.00 35.30 ATOM 1611 CG1 ILE A 210 127.009 83.101 14.378 1.00 36.13 ATOM 1612 CG2 ILE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1613 CD1 ILE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1614 N GLU A 211 123.135 81.303 13.271 1.00 34.62 ATOM 1615 CA GLU A 211 121.832 81.379 12.635 1.00 33.18 ATOM 1616 C GLU A 211 121.832 81.379 12.635 1.00 33.18 ATOM 1616 C GLU A 211 121.832 81.79 12.635 1.00 33.18 ATOM 1616 C GLU A 211 121.932 81.79 12.635 1.00 32.49 ATOM 1619 CG GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 122.657 79.924 10.751 1.00 34.14 ATOM 1620 CD GLU A 211 120.689 80.778 11.482 1.00 34.14 ATOM 1621 OEI GLU A 211 120.686 78.997 9.582 1.00 39.42 ATOM 1620 CD GLU A 211 120.686 78.997 9.582 1.00 39.42 ATOM 1621 OEI GLU A 211 120.967 79.924 10.751 1.00 36.42 ATOM 1622 OE2 GLU A 211 120.927 79.233 8.499 1.00 40.95 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1626 C PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1627 CB PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1628 C PHE A 212 119.913 77.673 13.807 1.00 27.07 ATOM 1629 CD1 PHE A 212 119.915 77.864 13.307 1.00 27.07 ATOM 1630 CD2 PHE A 212 119.957 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.867 77.920 11.955 1.00 27.46 ATOM 1634 N PRO A 213 120.928 80.375 20.991 1.00 30.49 ATOM 1636 C PRO A 213 120.928 80.375 20.991 1.00 30.49 ATOM 1637 O PRO A 213 120.928 80.375 20.991 1.00 30.49 ATOM 1636 C PRO A 213 120.928 80.375 20.991 1.00 30.49 ATOM 1636 C PRO A 213 120.928 80.375 20.991 1.00 32.42 ATOM 1637 O PRO A 213 120.938 81.168 11.755 1.00 27.46 ATOM 1640 C D PRO A 213 120.949 80.375 20.991 1.00 32.42 ATOM 1640 C D PRO A 213 120.949 80.375 20.991 1.00 32.42 ATOM 1640 C D PRO A 213 120.949 80.375 20.991 1.00 32.42 ATOM 1640 C D PRO A 213 120.949 80.375 20.991 1.00 32.42 ATOM 1640 C D PRO A 213 120.949 80.259														
ATOM 1610 CB LLE A 210 125.712 82.834 13.766 1.00 35.30 ATOM 1611 CG1 LLE A 210 127.090 83.101 14.378 1.00 36.13 ATOM 1612 CG2 LLE A 210 125.080 84.121 13.243 1.00 34.62 ATOM 1613 CD1 LLE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1614 N GLU A 211 121.832 81.179 12.635 1.00 33.34 ATOM 1615 CA GLU A 211 121.832 81.179 12.635 1.00 33.18 ATOM 1616 C GLU A 211 120.689 80.778 13.565 1.00 32.49 ATOM 1617 O GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.686 78.997 9.582 1.00 39.42 ATOM 1620 CD GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1621 OB1 GLU A 211 121.606 78.009 9.757 1.00 42.56 ATOM 1622 OEZ GLU A 211 120.952 79.893 14.526 1.00 33.18 ATOM 1624 CA PHE A 212 120.952 79.893 14.526 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1627 CB PHE A 212 119.187 77.673 13.807 1.00 27.07 ATOM 1629 CD1 PHE A 212 119.187 77.673 13.807 1.00 27.70 ATOM 1629 CD1 PHE A 212 119.187 77.673 13.807 1.00 27.70 ATOM 1630 CD2 PHE A 212 119.863 76.792 11.651 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.70 ATOM 1634 N PRO A 213 120.948 80.551 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.948 80.551 19.926 1.00 30.83 ATOM 1630 CD2 PHE A 212 119.863 76.792 11.651 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.70 ATOM 1634 N PRO A 213 120.948 80.551 19.926 1.00 30.59 ATOM 1635 CA PRO A 213 120.948 80.551 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.948 80.551 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.948 80.551 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.948 80.551 19.926 1.00 30.61 ATOM 1640 CD PRO A 213 120.958 82.663 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.958 80.551 19.926 1.00 30.61 ATOM 1640 CD PRO A 213 120.858 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.858 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.858 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.858 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.858 82	A	MOT	1608	С	ILE	Α	210	123.	357	82.142	14.291	1.00	33.53	C
ATOM 1611 CG1 ILE A 210 127.090 83.101 14.378 1.00 36.13 ATOM 1613 CD1 ILE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1613 CD1 ILE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1615 CA GLU A 211 123.135 81.320 13.271 1.00 33.34 ATOM 1616 C GLU A 211 120.689 80.778 13.565 1.00 32.49 ATOM 1616 C GLU A 211 120.689 80.778 13.565 1.00 32.49 ATOM 1617 O GLU A 211 119.572 81.266 13.405 1.00 32.49 ATOM 1618 CB GLU A 211 119.572 81.266 13.405 1.00 34.14 ATOM 1619 CG GLU A 211 120.687 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.687 79.924 10.751 1.00 36.42 ATOM 1621 OEI GLU A 211 120.687 79.924 10.751 1.00 36.42 ATOM 1621 OEI GLU A 211 120.668 78.997 9.582 1.00 39.42 ATOM 1622 OEZ GLU A 211 120.668 78.997 9.582 1.00 39.42 ATOM 1623 N PHE A 212 120.292 79.233 8.499 1.00 40.95 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1627 CB PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1628 CG PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1630 CD PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1631 CEI PHE A 212 119.510 77.864 13.307 1.00 27.41 ATOM 1630 CD PHE A 212 117.995 77.864 13.307 1.00 27.41 ATOM 1631 CEI PHE A 212 117.995 77.864 13.307 1.00 27.41 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 27.46 ATOM 1634 N PRO A 213 120.943 81.168 18.745 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.79 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.958 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.958 80.351 19.926 1.00 30.59 ATOM 1640 CD PRO A 213 120.958 80.531 19.926 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.865 82.	A	TOM	1609	0	ILE	A	210	122.	480	82.843	14.798	1.00	32.71	0
ATOM 1612 CG2 ILE A 210 125.080 84.121 13.243 1.00 34.62 ATOM 1613 CD1 ILE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1614 N GLU A 211 123.335 81.320 13.271 1.00 33.34 ATOM 1615 CA GLU A 211 121.832 81.179 12.635 1.00 33.18 ATOM 1616 C GLU A 211 121.832 81.179 12.635 1.00 33.18 ATOM 1617 O GLU A 211 121.832 81.179 12.635 1.00 33.49 ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.689 80.778 13.565 1.00 32.64 ATOM 1619 CG GLU A 211 120.667 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1621 OBI GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1622 OEZ GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 11.9912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.095 78.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD PHE A 212 119.187 77.673 13.807 1.00 27.70 ATOM 1630 CD2 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1630 CD2 PHE A 212 117.995 77.644 13.307 1.00 27.70 ATOM 1631 CEI PHE A 212 117.995 77.643 13.807 1.00 27.70 ATOM 1633 CZ PHE A 212 117.995 77.692 11.651 1.00 27.08 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.888 80.551 17.995 1.00 30.81 ATOM 1636 C PRO A 213 120.888 80.551 17.995 1.00 30.81 ATOM 1636 C PRO A 213 120.888 80.551 17.995 1.00 30.81 ATOM 1636 C PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1636 C PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766	A	TOM	1610	CB	ILE	Α	210	125.	712	82.834	13.766	1.00	35.30	C
ATOM 1613 CD1 ILE A 210 127.040 83.976 15.620 1.00 37.62 ATOM 1614 N GLU A 211 121.832 81.327 11.07 13.318 ATOM 1616 C GLU A 211 121.832 81.179 12.635 1.00 33.18 ATOM 1617 O GLU A 211 120.689 80.778 13.565 1.00 32.49 ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.657 79.924 10.751 1.00 34.14 ATOM 1620 CD GLU A 211 120.658 78.997 9.582 1.00 39.42 ATOM 1621 OE1 GLU A 211 120.658 78.997 9.582 1.00 39.42 ATOM 1622 CB2 GLU A 211 120.658 78.997 9.582 1.00 39.42 ATOM 1623 N PHE A 212 120.292 79.233 8.499 1.00 40.95 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 31.88 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.57 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1629 CD1 PHE A 212 119.912 79.465 13.307 1.00 27.87 ATOM 1629 CD1 PHE A 212 119.913 79.465 13.307 1.00 27.87 ATOM 1630 CD2 PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1631 CEI PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1631 CEI PHE A 212 117.905 77.864 13.307 1.00 27.87 ATOM 1630 CD2 PHE A 212 117.595 77.520 11.985 1.00 30.79 ATOM 1631 CEI PHE A 212 117.595 77.520 11.985 1.00 27.41 ATOM 1633 CZ PHE A 212 118.578 76.992 11.159 1.00 27.46 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1637 O PRO A 213 120.958 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.958 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.958 82.693 18.766 1.00 30.59 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1644 O HIS A 214 118.943 79.599 20.872 1.00 29.85 ATOM 1645 CB HIS A 214 117.993 79.599 20.872 1.00 29.85 ATOM 1646 CG HIS A 214 117.993 79.599 20.872 1.00 32.85 ATOM 1640 CD PRO A 213 119.606 82.949 17.659 1.00 30.59 ATOM 1640 CD PRO A 213 119.606 83.759 20.999 1.00 32.85 ATOM 1640 CD HIS A 214 116.896 82.663 21.9	A	MOT	1611	CG1	ILE	Α	210	127.	090	83.101	14.378	1.00	36.13	C
ATOM 1614 N GLU A 211 123.135 81.320 13.271 1.00 33.34 ATOM 1615 CA GLU A 211 121.832 81.179 12.635 1.00 33.34 ATOM 1616 C GLU A 211 120.689 80.778 13.565 1.00 32.49 ATOM 1617 O GLU A 211 120.689 80.778 13.565 1.00 32.49 ATOM 1617 O GLU A 211 120.689 80.778 13.565 1.00 32.64 ATOM 1618 CB GLU A 211 120.689 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.6657 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.666 78.009 9.757 1.00 36.42 ATOM 1622 OZZ GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1622 OZZ GLU A 211 120.952 79.893 14.526 1.00 30.657 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 30.657 ATOM 1624 CA PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1627 CB PHE A 212 120.705 78.622 17.578 1.00 29.04 ATOM 1628 CG PHE A 212 119.510 78.607 15.234 1.00 29.04 ATOM 1630 CDZ PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1630 CDZ PHE A 212 119.187 77.673 13.807 1.00 27.07 ATOM 1631 CEZ PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1631 CEZ PHE A 212 119.560 77.864 13.307 1.00 27.07 ATOM 1632 CZZ PHE A 212 119.567 77.520 11.985 1.00 27.08 ATOM 1632 CZZ PHE A 212 119.567 77.520 11.985 1.00 27.08 ATOM 1632 CZZ PHE A 212 119.567 77.520 11.985 1.00 27.08 ATOM 1632 CZZ PHE A 212 118.578 76.992 11.159 1.00 27.46 ATOM 1632 CZZ PHE A 212 118.578 76.992 11.159 1.00 27.46 ATOM 1633 CZ PHE A 212 118.578 76.992 11.159 1.00 27.46 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 CB PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1640 CD PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1640 CD PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1640 CD PRO A 213 1	A	MOT	1612	CG2	ILE	Α	210	125.	080	84.121	13.243	1.00	34.62	C
ATOM 1615 CA GLU A 211 121.832 81.179 12.635 1.00 33.18 ATOM 1616 C GLU A 211 120.689 80.778 13.565 1.00 32.64 ATOM 1617 O GLU A 211 121.943 80.186 11.482 1.00 32.64 ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1621 OE1 GLU A 211 120.666 78.099 9.757 1.00 42.56 ATOM 1622 OE2 GLU A 211 120.2968 78.8997 9.582 1.00 39.42 ATOM 1622 OE2 GLU A 211 120.2952 79.893 14.526 1.00 31.88 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 120.952 79.893 14.526 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.57 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1629 CD1 PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1629 CD1 PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1630 CD2 PHE A 212 117.595 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 117.595 77.864 13.307 1.00 27.70 ATOM 1633 CZ PHE A 212 117.595 77.864 13.307 1.00 27.70 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.948 80.531 19.926 1.00 30.59 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.79 ATOM 1636 C PRO A 213 120.948 80.531 19.926 1.00 30.40 29.65 ATOM 1636 C PRO A 213 120.948 80.531 19.926 1.00 30.40 29.65 ATOM 1636 C PRO A 213 120.988 80.531 19.926 1.00 30.42 ATOM 1636 C PRO A 213 120.988 80.531 19.926 1.00 30.42 ATOM 1637 O PRO A 213 120.988 80.531 19.926 1.00 30.42 ATOM 1640 CD PRO A 213 120.988 80.531 19.926 1.00 30.42 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.985 82.693 18.766 1.00 30.853 ATOM 1640 CD PRO A 2	A	MOT	1613	CD1	ILE	Α	210	127.	040	83.976	15.620	1.00	37.62	C
ATOM 1615 CA GLU A 211 121.832 81.179 12.635 1.00 33.18 ATOM 1616 C GLU A 211 120.689 80.778 13.565 1.00 32.64 ATOM 1617 O GLU A 211 119.572 81.266 13.405 1.00 32.64 ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1620 CD GLU A 211 120.686 78.997 9.582 1.00 39.42 ATOM 1621 OE1 GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1622 OE2 GLU A 211 120.868 78.997 9.757 1.00 42.56 ATOM 1622 OE2 GLU A 211 120.965 79.893 14.526 1.00 30.475 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1626 OPHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1629 CD1 PHE A 212 119.187 77.673 13.807 1.00 27.08 ATOM 1630 CD2 PHE A 212 117.995 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 117.995 77.864 13.307 1.00 27.70 ATOM 1633 CZ PHE A 212 117.995 77.864 13.307 1.00 27.70 ATOM 1633 CZ PHE A 212 117.995 77.864 13.307 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.79 ATOM 1636 C PRO A 213 120.948 80.531 19.926 1.00 30.59 ATOM 1638 CB PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1639 CP PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1630 CD PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1631 CB PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1634 N PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1635 CA PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1636 C PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1637 O PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1638 CB PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1640 CD PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1640 CD PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1640 CD PRO A 213 120.948 80.531 19.926 1.00 30.79 ATOM 1640 CD PRO A 213 120.958 82.663 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.958 82.663 19.959 1.00 32.42 ATOM 1640 CD PRO A 213 120.958 82.663 19.959 1.00 32.85 ATOM 1640 CD PRO A 213 120.958 82.663				N	GLU	Α	211	123.	135	81.320	13.271	1.00	33.34	N
ATOM 1617 O GLU A 211 119.572 81.266 13.405 1.00 32.64 ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1621 OE1 GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1622 OE2 GLU A 211 120.925 79.233 8.499 1.00 40.95 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1626 O PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 119.510 78.607 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 119.187 77.673 13.807 1.00 27.41 ATOM 1630 CD2 PHE A 212 119.963 76.792 11.651 1.00 27.70 ATOM 1631 CB1 PHE A 212 117.995 77.864 13.307 1.00 27.70 ATOM 1633 CZ PHE A 212 117.995 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1635 CA PRO A 213 120.493 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1638 CB PRO A 213 120.943 80.850 17.396 1.00 30.59 ATOM 1639 CG PRO A 213 120.928 80.850 17.396 1.00 30.59 ATOM 1630 CD PRO A 213 120.928 80.850 17.396 1.00 30.59 ATOM 1631 CB PRO A 213 120.928 80.850 17.396 1.00 30.59 ATOM 1634 N PRO A 213 120.928 80.850 17.396 1.00 30.59 ATOM 1636 C PRO A 213 120.928 80.850 17.396 1.00 30.59 ATOM 1637 O PRO A 213 120.928 80.551 19.926 1.00 30.81 ATOM 1640 CD PRO A 213 120.928 80.375 20.991 1.00 22.06 ATOM 1640 CD PRO A 213 120.928 80.375 20.991 1.00 32.42 ATOM 1646 CG PRO A 213 120.928 80.375 20.991 1.00 32.42 ATOM 1646 CG PRO A 213 120.828 80.350 19.926 1.00 30.81 ATOM 1640 CD PRO A 213 120.929 80.375 20.991 1.00 32.42 ATOM 1644 O HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1646 CG PRO A 213 110.017.93 81.999 1.00 26.06 ATOM 1640 CD PRO A 213 110.017.93 81.999 1.00 26.06 ATOM 1640 CD PRO A 213 110.017.94 81.999 1.00 37.66 ATOM 1646 CB HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1646 CD HIS A 214 116.854 82.469 19.776 1.00	A	MOT	1615	CA	GLU	Α	211	121.	832	81.179	12.635	1.00	33.18	C
ATOM 1618 CB GLU A 211 121.943 80.186 11.482 1.00 34.14 ATOM 1619 CG GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.668 78.997 9.582 1.00 39.42 ATOM 1621 OE1 GLU A 211 120.668 78.997 9.582 1.00 39.42 ATOM 1622 OE2 GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 29.04 ATOM 1627 CB PHE A 212 119.510 78.007 15.224 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1630 CD2 PHE A 212 119.912 79.861 10.00 27.08 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1633 CZ PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 27.46 ATOM 1635 CZ PHE A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.851 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.851 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1637 O PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1637 O PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1638 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1640 C PRO A 213 120.208 80.531 19.926 1.00 30.81 ATOM 1640 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1640 C PRO A 213 120.948 80.531 19.926 1.00 30.59 ATOM 1640 C PRO A 213 120.948 80.531 19.926 1.00 30.59 ATOM 1640 C PRO A 213 120.949 80.375 20.991 1.00 32.42 ATOM 1640 C PRO A 213 120.949 80.375 20.991 1.00 32.42 ATOM 1640 C PRO A 213 120.949 80.375 20.991 1.00 30.59 ATOM 1640 C PRO A 213 120.949 90.375 1.00 30.59 ATOM 1640 C PRO A 213 120.949 90.375 1.00 30.59 ATOM 1640 C PRO A	A	TOM	1616	C	GLU	Α	211	120.	689	80.778	13.565	1.00	32.49	C
ATOM 1619 CG GLU A 211 120.657 79.924 10.751 1.00 36.42 ATOM 1620 CD GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1621 OE1 GLU A 211 121.606 78.009 9.757 1.00 42.56 ATOM 1622 OE2 GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1622 OE2 GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 27.87 ATOM 1628 CG PHE A 212 119.510 78.007 15.234 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1633 CZ PHE A 212 119.586 76.792 11.651 1.00 27.08 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1638 CB PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.958 80.531 19.926 1.00 30.81 ATOM 1636 C PRO A 213 120.98 80.531 19.926 1.00 30.81 ATOM 1640 CD PRO A 213 120.993 80.375 20.991 1.00 32.42 ATOM 1640 CD PRO A 213 120.185 80.258 20.991 1.00 30.72 ATOM 1640 CD PRO A 213 120.186 82.949 17.850 1.00 30.59 ATOM 1640 CD PRO A 213 120.178 82.091 17.850 1.00 30.59 ATOM 1640 CD PRO A 213 120.178 82.091 17.850 1.00 30.59 ATOM 1640 CD PRO A 213 120.178 82.091 17.850 1.00 30.59 ATOM 1640 CD PRO A 213 120.178 82.091 17.00 30.72 ATOM 1640 CD PRO A 213 120.178 82.091 17.850 1.00	A	MOT	1617	0	GLU	Α	211	119.	572	81.266	13.405	1.00	32.64	0
ATOM 1620 CD GLU A 211 120.868 78.997 9.582 1.00 39.42 ATOM 1621 OE1 GLU A 211 121.606 78.009 9.757 1.00 42.56 ATOM 1622 OE2 GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 C PHE A 212 119.510 78.622 17.578 1.00 29.04 ATOM 1629 CD PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD PHE A 212 119.187 77.673 13.807 1.00 27.41 ATOM 1630 CD2 PHE A 212 119.187 77.673 13.807 1.00 27.41 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1632 CZ PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.928 80.531 19.926 1.00 30.81 ATOM 1637 O PRO A 213 120.893 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.893 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.893 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.893 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.855 82.099 1.00 32.42 ATOM 1640 CD PRO A 213 120.855 82.099 1.00 32.85 ATOM 1640 CD PRO A 213 120.855 82.099 1.00 30.79 AT	A	MOT	1618	CB	GLU	Α	211	121.	943	80.186	11.482	1.00	34.14	C
ATOM 1621 OE1 GLU A 211 121.606 78.009 9.757 1.00 42.56 ATOM 1622 OE2 GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1633 CZ PHE A 212 119.595 77.520 11.985 1.00 27.46 ATOM 1633 CZ PHE A 213 120.482 80.850 17.396 1.00 27.46 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1638 CB PRO A 213 120.208 80.531 19.926 1.00 30.81 ATOM 1639 CG PRO A 213 120.208 80.531 19.926 1.00 30.81 ATOM 1639 CG PRO A 213 120.882 82.693 18.765 1.00 32.42 ATOM 1638 CB PRO A 213 120.883 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.79 ATOM 1640 CD PRO A 213 120.853 82.693 10.00 20.793 1.00 28.54 ATOM 1640 CD PRO A 213 120.804 80.850 1.00 20.793 1.00 28.54 ATOM 1640 CD PRO A 2	A	TOM	1619	CG	GLU	A	211	120.	657	79.924	10.751	1.00	36.42	C
ATOM 1622 OE2 GLU A 211 120.292 79.233 8.499 1.00 40.95 ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1630 CD2 PHE A 212 119.863 76.792 11.651 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 117.595 77.520 11.985 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1637 O PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1639 CG PRO A 213 120.792 80.375 20.991 1.00 30.59 ATOM 1640 CD PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 17.983 78.090 20.793 1.00 29.65 ATOM 1640 CD PRO A 214 118.193 79.589 20.872 1.00 29.81 ATOM 1644 O HIS A 214 118.193 79.589 20.872 1.00 29.81 ATOM 1646 CG HIS A 214 116.876 81.752 20.955 1.00 36.67 ATOM 1647 NDI HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1648 CD2 HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1649 CB1 HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1649 CB1 HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.	A	MOT	1620	CD	GLU	A	211	120.	868	78.997	9.582	1.00	39.42	C
ATOM 1623 N PHE A 212 120.952 79.893 14.526 1.00 31.88 ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.70 ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.551 1.00 27.08 ATOM 1632 CE2 PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.59 ATOM 1638 CB PRO A 213 120.208 80.531 19.926 1.00 30.42 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1642 CA HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1643 C HIS A 214 118.943 79.589 20.872 1.00 29.81 ATOM 1643 C HIS A 214 118.943 79.589 20.872 1.00 29.81 ATOM 1644 O HIS A 214 117.983 78.090 20.793 1.00 28.54 ATOM 1646 CG HIS A 214 116.865 81.752 20.995 1.00 32.85 ATOM 1647 ND1 HIS A 214 116.876 81.752 20.995 1.00 32.85 ATOM 1648 CB HIS A 214 116.876 81.752 20.995 1.00 32.85 ATOM 1649 CB HIS A 214 116.876 81.752 20.995 1.00 33.852 ATOM 1649 CB HIS A 214 116.896 82.469 19.776 1.00 33.33 ATOM 1648 CD HIS A 214 116.896 82.469 19.776 1.00 38.92 ATOM 1649 CB HIS A 214 116.998 82.663 21.999 1.00 37.66 ATOM 1649 CB HIS A 214 116.998 82.663 21.999 1.00 37.66 ATOM 1649 CB HIS A 214 116.998 82.663 21.999 1.00 37.66 ATOM 1649 CB HIS A 214 116.876 81.752 20.955 1.00 38.52 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.999 1.00 37.66 ATOM 1649 CB HIS A 214 116.998 82.663 21.999 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.960 83.758 20.047 1.00 38.92 ATOM 1651 N ALA A 215 118.633 75.984 19.774 1.00 26.19	A	MOT	1621	OE1	GLU	A	211	121.	606	78.009	9.757	1.00	42.56	0
ATOM 1624 CA PHE A 212 119.912 79.465 15.466 1.00 30.57 ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 117.595 77.520 11.985 1.00 27.46 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.81 ATOM 1637 O PRO A 213 120.208 80.531 19.926 1.00 30.81 ATOM 1638 CB PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.72 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1644 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1645 CB HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1646 CG HIS A 214 117.983 78.090 20.793 1.00 29.85 ATOM 1646 CB HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1646 CB HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1646 CD HIS A 214 116.876 81.752 20.955 1.00 36.67 ATOM 1648 CD2 HIS A 214 116.898 82.663 21.949 1.00 38.52 ATOM 1649 CEI HIS A 214 116.998 82.663 21.949 1.00 38.52 ATOM 1649 CEI HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CEI HIS A 214 116.998 82.663 21.949 1.00 38.52 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1651 N ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19	A	TOM	1622	OE2	GLU	A	211	120.	292	79.233	8.499	1.00	40.95	0
ATOM 1625 C PHE A 212 120.409 79.610 16.902 1.00 30.83 ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 117.595 77.520 11.651 1.00 27.08 ATOM 1633 CZ PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 118.578 76.992 11.159 1.00 27.46 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1638 CB PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1642 CA HIS A 214 118.193 79.589 20.872 1.00 29.81 ATOM 1644 O HIS A 214 117.133 77.549 21.499 1.00 26.06 ATOM 1645 CB HIS A 214 116.876 81.752 20.995 1.00 36.67 ATOM 1646 CG HIS A 214 116.876 81.752 20.995 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.876 81.752 20.995 1.00 38.33 ATOM 1649 CE1 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CEI HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CEI HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 83.963 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1652 CA ALA A 215 118.633 75.984 19.777 1.00 26.19	A	MOT	1623	N	PHE	A	212	120.	952	79.893	14.526	1.00	31.88	N
ATOM 1626 O PHE A 212 120.705 78.622 17.578 1.00 30.70 ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1637 O PRO A 213 120.828 80.531 19.926 1.00 30.81 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 29.65 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 30.72 ATOM 1643 C HIS A 214 118.944 80.156 19.757 1.00 30.72 ATOM 1644 O HIS A 214 118.944 80.156 19.757 1.00 29.81 ATOM 1645 CB HIS A 214 117.983 78.090 20.793 1.00 29.81 ATOM 1646 CG HIS A 214 117.983 78.090 20.793 1.00 28.54 ATOM 1646 CG HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1647 ND1 HIS A 214 116.854 82.469 19.776 1.00 30.66 ATOM 1648 CD2 HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1649 CE1 HIS A 214 116.854 82.469 19.776 1.00 38.53 ATOM 1649 CE1 HIS A 214 116.854 82.469 19.776 1.00 38.52 ATOM 1650 NE2 HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.898 82.663 21.949 1.00 37.66 ATOM 1651 N ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOT	1624	CA	PHE	A	212	119.	912	79.465	15.466	1.00	30.57	C
ATOM 1627 CB PHE A 212 119.510 78.007 15.234 1.00 29.04 ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1637 O PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1638 CB PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.868 82.949 17.850 1.00 29.65 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1642 CA HIS A 214 117.983 78.090 20.872 1.00 29.81 ATOM 1644 O HIS A 214 117.983 78.090 20.793 1.00 28.54 ATOM 1646 CG HIS A 214 117.983 78.090 20.793 1.00 28.54 ATOM 1646 CG HIS A 214 116.856 81.752 20.995 1.00 36.67 ATOM 1646 CG HIS A 214 116.856 81.752 20.995 1.00 37.66 ATOM 1646 CD HIS A 214 116.856 81.752 20.995 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1649 CE1 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1651 N ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOT	1625	C	PHE	Α	212	120.	409	79.610	16.902		30.83	C
ATOM 1628 CG PHE A 212 119.187 77.673 13.807 1.00 27.87 ATOM 1629 CD1 PHE A 212 120.161 77.139 12.969 1.00 27.41 ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1637 O PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1638 CB PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1642 CA HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1643 C HIS A 214 118.193 79.589 20.872 1.00 29.81 ATOM 1644 O HIS A 214 117.983 78.090 20.793 1.00 28.54 ATOM 1646 CG HIS A 214 116.876 81.752 20.995 1.00 32.85 ATOM 1646 CG HIS A 214 116.876 81.752 20.995 1.00 32.85 ATOM 1646 CG HIS A 214 116.876 81.752 20.955 1.00 32.85 ATOM 1648 CD2 HIS A 214 116.876 81.752 20.955 1.00 36.67 ATOM 1648 CD2 HIS A 214 116.876 81.752 20.955 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.894 82.469 19.776 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.896 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.896 82.663 21.949 1.00 37.66 ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1650 NE2 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 116.960 83.758 20.047 1.00 38.92 ATOM 1650 NE2 HIS A 214 116.960 83.758 20.047 1.00 38.92 ATOM 1650 NE2 HIS A 214 116.834 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.603 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.603 75.263 21.100 1.00 25.46	A	TOM	1626	0										0
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ATOM 1630 CD2 PHE A 212 117.905 77.864 13.307 1.00 27.70 ATOM 1631 CE1 PHE A 212 119.863 76.792 11.651 1.00 27.08 ATOM 1632 CE2 PHE A 212 117.595 77.520 11.985 1.00 28.23 ATOM 1633 CZ PHE A 212 118.578 76.982 11.159 1.00 27.46 ATOM 1634 N PRO A 213 120.482 80.850 17.396 1.00 30.79 ATOM 1635 CA PRO A 213 120.943 81.168 18.745 1.00 30.59 ATOM 1636 C PRO A 213 120.208 80.531 19.926 1.00 30.81 ATOM 1637 O PRO A 213 120.792 80.375 20.991 1.00 32.42 ATOM 1638 CB PRO A 213 120.853 82.693 18.766 1.00 30.59 ATOM 1639 CG PRO A 213 119.680 82.949 17.850 1.00 29.65 ATOM 1640 CD PRO A 213 120.117 82.091 16.697 1.00 30.72 ATOM 1641 N HIS A 214 118.944 80.156 19.757 1.00 31.01 ATOM 1642 CA HIS A 214 118.193 79.589 20.872 1.00 29.81 ATOM 1644 O HIS A 214 117.983 78.090 20.793 1.00 28.54 ATOM 1645 CB HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1646 CG HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1646 CD HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1646 CG HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1648 CD2 HIS A 214 116.896 81.752 20.955 1.00 36.67 ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.998 82.663 21.949 1.00 38.52 ATOM 1650 NE2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1651 N ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19														C
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ATOM 1645 CB HIS A 214 116.825 80.258 20.990 1.00 32.85 ATOM 1646 CG HIS A 214 116.876 81.752 20.955 1.00 36.67 ATOM 1647 ND1 HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 117.049 83.903 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOT	1643	С	HIS	Α	214	117.	983	78.090	20.793	1.00	28.54	C
ATOM 1646 CG HIS A 214 116.876 81.752 20.955 1.00 36.67 ATOM 1647 ND1 HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 117.049 83.903 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOTA	1644	0	HIS	A	214	117.	133	77.549	21.499	1.00	26.06	0
ATOM 1647 ND1 HIS A 214 116.854 82.469 19.776 1.00 38.33 ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 117.049 83.903 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOT	1645	CB	HIS	Α	214	116.	825	80.258	20.990	1.00	32.85	C
ATOM 1648 CD2 HIS A 214 116.998 82.663 21.949 1.00 37.66 ATOM 1649 CE1 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 117.049 83.903 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOT	1646	CG	HIS	A	214	116.	876	81.752	20.955	1.00	36.67	C
ATOM 1649 CE1 HIS A 214 116.960 83.758 20.047 1.00 38.52 ATOM 1650 NE2 HIS A 214 117.049 83.903 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOT	1647	ND1	HIS	A	214	116.	854	82.469	19.776	1.00	38.33	N
ATOM 1650 NE2 HIS A 214 117.049 83.903 21.357 1.00 38.92 ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOTA	1648	CD2	HIS	A	214	116.	998	82.663	21.949	1.00	37.66	C
ATOM 1651 N ALA A 215 118.756 77.424 19.943 1.00 27.04 ATOM 1652 CA ALA A 215 118.633 75.984 19.774 1.00 26.19 ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46	A	MOTA	1649	CE1	HIS	A	214							C
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ATOM 1653 C ALA A 215 118.800 75.263 21.100 1.00 25.46														N
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ATOM 1654 O ALA A 215 119.618 /5.658 21.924 1.00 25.60														C
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ATOM	1655	CB	ALA	Α	215	119.667	75.486	18.768	1.00	25.12	C
ATOM	1656	N	ALA	A	216	118.004	74.216	21.305	1.00		N
ATOM	1657	CA	ALA	A	216	118.072	73.417	22.523		25.69	C
MOTA	1658	C	ALA	A	216	119.095	72.341	22.211		26.07	C
MOTA	1659	0	ALA	A	216	118.880	71.516	21.327	1.00	27.31	0
MOTA	1660	CB	ALA	A	216	116.714	72.797	22.824	1.00	25.56	C
MOTA	1661	N	LYS	A	217	120.212	72.355	22.931	1.00	26.34	N
MOTA	1662	CA	LYS	A	217	121.289	71.409	22.677	1.00	25.73	С
MOTA	1663	C	LYS	A	217	121.786	70.689	23.918	1.00	26.81	C
ATOM	1664	0	LYS	A	217	121.480	71.084	25.044	1.00		0
MOTA	1665	CB	LYS	Α	217	122.450	72.150	22.015		25.06	C
MOTA	1666	CG	LYS	A	217	122.081	72.771	20.672		24.94	C
MOTA	1667	CD	LYS	Α	217	123.162	73.693	20.150		24.62	C
ATOM	1668	CE	LYS	Α	217	122.830	74.159	18.749	1.00		C
ATOM	1669	NZ	LYS	A	217	123.873	75.063	18.196		28.65	N
ATOM	1670	N	ASP	Α	218	122.546	69.616	23.700	1.00		N
ATOM	1671	CA	ASP	A	218	123.106	68.855	24.808		29.53	C
ATOM	1672	C			218	124.458	69.434	25.200	1.00		C
ATOM	1673	0	ASP	A	218	124.864	70.494	24.715		27.51	0
ATOM	1674	CB	ASP			123.270	67.363	24.458	1.00		C
ATOM	1675	CG	ASP			124.142	67.130	23.231		32.50	C
ATOM	1676					125.144	67.853	23.060		33.09	0
ATOM	1677	OD2	ASP			123.848	66.200	22.447	1.00		0
ATOM	1678	N	GLU			125.143	68.718	26.082	1.00		N
ATOM	1679	CA	GLU			126.445	69.117	26.586		33.04	C
ATOM	1680	C	GLU			127.492	69.306	25.505		31.97	C
ATOM	1681	0	GLU			128.455	70.037	25.697	1.00		0
MOTA	1682	CB	GLU			126.931	68.087	27.613		36.51	C
ATOM	1683	CG	GLU			126.807	66.636	27.148		43.30	C
ATOM	1684	CD	GLU			127.129	65.626	28.245		46.43	C
ATOM	1685	OE1	GLU			126.570	65.740	29.358		49.99	0
ATOM	1686	OE2	GLU			127.926	64.702	27.994		49.56	
ATOM	1687	N			220	127.297	68.663	24.363		31.08	N C
ATOM	1688	CA			220	128.253	68.753	23.264		30.41	C
ATOM	1689	C			220	127.863	69.773	22.205		30.02 30.29	0
ATOM	1690	O			220	128.534	69.897 67.378	21.180 22.619		30.42	C
ATOM	1691	CB			220	128.401	66.292	23.600		31.61	C
ATOM	1692	CG	PHE PHE		220	128.737 129.949	66.306	24.287	1.00		C
ATOM	1693	CD1 CD2			220 220	127.831	65.267	23.855	1.00		C
ATOM	1694	CE1			220	130.252	65.317	25.209	1.00		C
ATOM ATOM	1695 1696		PHE			128.124	64.276	24.775	1.00		C
ATOM	1697	CEZ			220	129.337	64.299	25.453		31.56	C
ATOM	1698	N			221	126.773	70.496	22.453		30.25	N
ATOM	1699	CA			221	126.773	71.485	21.498		29.06	C
ATOM	1700	C			221	125.573	70.852	20.325		28.96	C
ATOM	1701	0			221	125.575	71.417	19.232		27.40	0
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ATOM	1702	N	ARG	A	222	125.029	69.658	20.551	1.00	28.41	N
ATOM	1703	CA	ARG	Α	222	124.269	68.949	19.534	1.00	27.01	С
ATOM	1704	С	ARG	Α	222	122.797	69.105	19.909	1.00	25.67	C
ATOM	1705	0	ARG	Α	222	122.448	69.100	21.081	1.00	24.90	0
ATOM	1706	CB	ARG	Α	222	124.640	67.470	19.518	1.00	26.66	C
ATOM	1707	CG	ARG	Α	222	126.102	67.212	19.270	1.00	27.12	C
ATOM	1708	CD	ARG	A	222	126.398	65.719	19.306	1.00	28.42	C
ATOM	1709	NE	ARG	Α	222	125.961	65.097	20.559	1.00	30.08	N
ATOM	1710	CZ	ARG	Α	222	126.067	63.799	20.827	1.00	27.75	C
ATOM	1711	NH1	ARG	Α	222	126.599	62.973	19.934	1.00	25.60	N
ATOM	1712	NH2	ARG	Α	222	125.620	63.328	21.981	1.00	27.54	N
ATOM	1713	N	LEU	A	223	121.944	69.248	18.906	1.00	25.37	N
ATOM	1714	CA	LEU	Α	223	120.506	69.417	19.129	1.00	25.09	C
ATOM	1715	С	LEU	Α	223	119.943	68.277	19.967	1.00	25.23	C
ATOM	1716	0	LEU	Α	223	120.390	67.128	19.853	1.00	25.90	0
ATOM	1717	CB	LEU	A	223	119.766	69.464	17.782	1.00	22.94	C
ATOM	1718	CG	LEU	Α	223	120.155	70.538	16.755	1.00	23.39	C
ATOM	1719	CD1	LEU	Α	223	119.455	70.265	15.420	1.00	22.13	C
ATOM	1720	CD2	LEU	Α	223	119.802	71.917	17.288	1.00	22.20	C
ATOM	1721	N	LEU	Α	224	118.963	68.595	20.811	1.00	24.72	N
ATOM	1722	CA	LEU	A	224	118.318	67.585	21.647	1.00	24.86	C
ATOM	1723	C	LEU	A	224	117.331	66.806	20.806	1.00	24.46	C
ATOM	1724	0	LEU	A	224	116.871	67.281	19.775	1.00	24.23	0
ATOM	1725	CB	LEU	A	224	117.545	68.226	22.798	1.00	23.92	C
ATOM	1726	CG	LEU	A	224	118.313	68.961	23.888	1.00	25.09	C
MOTA	1727	CD1	LEU	A	224	117.344	69.527	24.924	1.00	21.77	C
ATOM	1728	CD2	LEU	A	224	119.295	67.987	24.529	1.00	25.24	C
ATOM	1729	N	VAL	Α	225	117.008	65.600	21.243	1.00	25.32	N
ATOM	1730	CA	VAL	Α	225	116.027	64.803	20.532	1.00	25.61	C
MOTA	1731	C	VAL	A	225	115.498	63.714	21.440	1.00	25.77	C
ATOM	1732	0	VAL	Α	225	116.227	63.181	22.282	1.00	25.12	0
ATOM	1733	CB	VAL	Α	225	116.601	64.151	19.237	1.00	27.42	C
ATOM	1734	CG1	VAL	Α	225	117.632	63.069	19.577	1.00	26.05	C
ATOM	1735	CG2	VAL	Α	225	115.456	63.574	18.400	1.00	26.21	C
ATOM	1736	N	ALA	A	226	114.214	63.412	21.272	1.00	24.98	N
ATOM	1737	CA	ALA	Α	226	113.547	62.373	22.037	1.00	23.32	C
ATOM	1738	C	ALA	Α	226	112.985	61.398	21.013	1.00	23.19	C
ATOM	1739	0	ALA	Α	226	112.839	61.734	19.838	1.00	22.48	0
ATOM	1740	CB	ALA	A	226	112.426	62.966	22.876	1.00	22.70	C
ATOM	1741	N	ALA	A	227	112.692	60.184	21.452	1.00	22.90	N
ATOM	1742	CA	ALA	Α	227	112.143	59.179	20.560	1.00	24.67	C
ATOM	1743	C	ALA			111.116	58.381	21.346	1.00	26.09	C
ATOM	1744	0	ALA	A	227	111.237	58.219	22.566	1.00	26.01	0
ATOM	1745	CB	ALA			113.255	58.266	20.029		22.84	C
ATOM	1746	N	ALA			110.098	57.892	20.650		26.88	N
ATOM	1747	CA	ALA			109.053	57.130	21.306		27.92	C
ATOM	1748	C	ALA	A	228	109.201	55.630	21.125	1.00	29.27	C

ATOM	1749	0	ALA	A	228	109.788	55.149	20.153	1.00	28.65	0
ATOM	1750	CB	ALA	Α	228	107.697	57.575	20.792	1.00	28.14	C
ATOM	1751	N	VAL	Α	229	108.662	54.901	22.091	1.00	30.70	N
ATOM	1752	CA	VAL	Α	229	108.654	53.447	22.082	1.00	31.77	C
ATOM	1753	C	VAL	Α	229	107.354	53.075	22.785	1.00	33.71	C
MOTA	1754	0	VAL	Α	229	106.778	53.897	23.508	1.00	34.20	0
MOTA	1755	CB	VAL	Α	229	109.847	52.850	22.876	1.00	31.31	C
ATOM	1756	CG1	VAL	A	229	111.156	53.346	22.298	1.00	31.39	C
MOTA	1757	CG2	VAL	A	229	109.739	53.208	24.357	1.00	31.01	C
ATOM	1758	N	GLY	A	230	106.871	51.858	22.570	1.00	34.88	N
ATOM	1759	CA	GLY	Α	230	105.652	51.453	23.240	1.00	35.58	C
ATOM	1760	C	GLY	Α	230	105.976	50.476	24.352	1.00	36.88	C
ATOM	1761	0	GLY	Α	230	107.100	50.411	24.840	1.00	36.39	0
ATOM	1762	N	VAL	Α	231	104.975	49.722	24.772	1.00	39.52	N
MOTA	1763	CA	VAL	Α	231	105.176	48.719	25.801	1.00	41.92	C
ATOM	1764	C	VAL	Α	231	104.905	47.363	25.140	1.00	43.78	C
ATOM	1765	0	VAL	Α	231	103.779	46.858	25.143	1.00	44.73	0
ATOM	1766	CB	VAL	Α	231	104.240	48.966	26.997	1.00	41.08	C
ATOM	1767	CG1	VAL	Α	231	104.639	50.257	27.691	1.00	40.96	C
ATOM	1768	CG2	VAL	Α	231	102.810	49.071	26.526	1.00	40.96	C
ATOM	1769	N	THR	Α	232	105.954	46.807	24.538	1.00	44.60	N
ATOM	1770	CA	THR	Α	232	105.881	45.532	23.845	1.00	45.87	C
ATOM	1771	C	THR	A	232	107.106	44.710	24.200	1.00	46.68	C
ATOM	1772	0	THR	A	232	107.838	45.059	25.120	1.00	47.51	0
ATOM	1773	CB	THR	A	232	105.842	45.739	22.328	1.00	46.37	C
ATOM	1774	OG1	THR	A	232	107.038	46.404	21.906	1.00	46.75	0
ATOM	1775	CG2	THR	Α	232	104.645	46.590	21.946	1.00	46.43	C
ATOM	1776	N	SER	Α	233	107.342	43.626	23.468	1.00	48.35	N
MOTA	1777	CA	SER	Α	233	108.491 -	42.768	23.755	1.00	49.99	C
ATOM	1778	C	SER	Α	233	109.804	43.315	23.199	1.00	50.98	C
MOTA	1779	0	SER	Α	233	110.887	42.949	23.667	1.00	52.13	0
ATOM	1780	CB	SER	Α	233	108.256	41.355	23.211	1.00	49.80	C
ATOM	1781	OG	SER	A	233	108.089	41.370	21.809	1.00	51.16	0
MOTA	1782	N	ASP	Α	234	109.710	44.197	22.210	1.00	51.06	N
ATOM	1783	CA	ASP	A	234	110.901	44.779	21.606	1.00	51.16	C
MOTA	1784	С	ASP	A	234	111.207	46.174	22.142	1.00	50.20	C
ATOM	1785	0	ASP	Α	234	112.024	46.892	21.570	1.00	50.68	0
ATOM	1786	CB	ASP	A	234	110.733	44.835	20.092	1.00	52.65	C
MOTA	1787	CG	ASP	Α	234	109.474	45.563	19.686	1.00	55.59	C
MOTA	1788	OD1	ASP	Α	234	108.388	45.134	20.138	1.00	57.24	0
ATOM	1789	OD2	ASP	A	234	109.566	46.556	18.924	1.00	56.62	0
MOTA	1790	N	THR	Α	235	110.559	46.553	23.239		48.92	N
MOTA	1791	CA	THR	Α	235	110.775	47.866	23.837		48.36	C
MOTA	1792	C	THR	Α	235	112.219	48.115	24.279		48.23	C
MOTA	1793	0	THR	A	235	112.835	49.092	23.855		48.06	0
MOTA	1794	CB			235	109.844	48.080	25.035		48.22	C
ATOM	1795	OG1	THR	A	235	108.492	48.121	24.572	1.00	48.44	0

ATOM	1796	CG2	THR	A	235	110.164	49.382	25.740	1.00 48.44	С
MOTA	1797	N	PHE	A	236	112.755	47.248	25.138	1.00 47.72	N
MOTA	1798	CA	PHE	A	236	114.133	47.403	25.594	1.00 46.43	C
MOTA	1799	C	PHE	A	236	115.075	47.536	24.420	1.00 45.57	C
MOTA	1800	0	PHE	A	236	115.856	48.480	24.342	1.00 45.91	0
MOTA	1801	CB	PHE	A	236	114.568	46.209	26.434	1.00 47.23	C
MOTA	1802	CG	PHE	A	236	114.351	46.394	27.892	1.00 48.55	C
MOTA	1803	CD1	PHE	A	236	115.014	47.402	28.572	1.00 49.39	C
ATOM	1804	CD2	PHE	Α	236	113.483	45.567	28.591	1.00 50.16	C
MOTA	1805	CE1	PHE	Α	236	114.817	47.588	29.931	1.00 51.05	C
ATOM	1806	CE2	PHE	Α	236	113.276	45.742	29.959	1.00 51.03	C
ATOM	1807	CZ			236	113.945	46.755	30.630	1.00 51.13	C
ATOM	1808	N	GLU			114.998	46.570	23.513	1.00 44.78	N
ATOM	1809	CA	GLU			115.835	46.548	22.320	1.00 44.06	C
ATOM	1810	C	GLU		237	115.749	47.901	21.626	1.00 41.89	C
MOTA	1811	0	GLU			116.756	48.466	21.210	1.00 41.21	0
ATOM	1812	CB	GLU			115.334	45.461	21.375	1.00 46.63	C
ATOM	1813	CG			237	116.258	45.132	20.226	1.00 50.92	C
ATOM	1814	CD	GLU			115.598	44.192	19.233	1.00 54.10	C
ATOM	1815	OE1			237	114.939	43.224	19.686	1.00 55.04	0
ATOM	1816	OE2	GLU			115.749	44.410	18.004	1.00 55.45	0
ATOM	1817	N	ARG			114.524	48.410	21.521	1.00 40.47	N
ATOM	1818	CA			238	114.244	49.691	20.880	1.00 39.30	C
ATOM	1819	C			238	114.820	50.864	21.667	1.00 38.27	C
ATOM	1820	O	ARG			115.508	51.715	21.109	1.00 37.62	0
ATOM	1821	CB	ARG			112.738	49.888	20.743	1.00 38.03	C
ATOM	1822	CG			238	112.380 110.901	50.858	19.656 19.586	1.00 38.82 1.00 39.86	C
ATOM	1823	CD NE	ARG ARG		238	110.555	51.126 51.547	18.236	1.00 33.88	N
ATOM	1824	CZ			238	110.333	50.707	17.233	1.00 42.02	C
ATOM	1825 1826	NH1			238	110.332	49.402	17.430	1.00 44.36	N
ATOM ATOM	1827	NH2	ARG			110.410	51.170	16.025	1.00 43.46	N
ATOM	1828	N	ALA			114.518	50.904	22.962	1.00 43.40	N
ATOM	1829	CA	ALA			114.991	51.960	23.848	1.00 37.99	C
MOTA	1830	C	ALA			116.516	52.030	23.869	1.00 37.33	C
ATOM	1831	0	ALA			117.093	53.121	23.815	1.00 39.71	0
ATOM	1832	СВ	ALA			114.461	51.729	25.249	1.00 36.95	C
ATOM	1833	N	GLU			117.174	50.873	23.944	1.00 41.10	N
ATOM	1834	CA	GLU			118.631	50.852	23.965	1.00 41.70	C
ATOM	1835	C	GLU			119.239	51.237	22.621	1.00 40.70	C
ATOM	1836	0	GLU			120.358	51.744	22.570	1.00 40.79	0
ATOM	1837	CB	GLU			119.157	49.486	24.428	1.00 44.05	C
ATOM	1838	CG	GLU			118.629	48.286	23.678	1.00 48.07	C
ATOM	1839	CD	GLU			119.141	46.973	24.266	1.00 51.41	C
ATOM	1840		GLU			119.010	46.776	25.499	1.00 51.93	0
ATOM	1841					119.664	46.135	23.496	1.00 52.53	0
ATOM	1842	N	ALA			118.508	51.005	21.536	1.00 39.57	N

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### ALA A 241 118.999 51.373 20.208 MOTA 1843 CA 1.00 39.09 MOTA **ALA A 241** 1844 C 118.911 52.893 20.071 1.00 38.48 119.805 ATOM 1845 ALA A 241 0 53.533 19.521 1.00 38.30 1846 **ALA A 241** MOTA CB 118.161 50.701 19.125 1.00 39.36 ATOM LEU A 242 1847 N 117.822 53.460 20.579 1.00 37.48 **LEU A 242** 117.603 20.549 1.00 36.64 ATOM 1848 CA 54.902 **LEU A 242** ATOM 1849 55.648 C 21.421 118.607 1.00 36.34 **LEU A 242** 119.217 **ATOM** 1850 56.620 1.00 35.49 0 20.973 **LEU A 242** MOTA 1851 CB 116.177 55.212 1.00 36.24 20.999 1852 CG LEU A 242 55.051 1.00 37.63 MOTA 115.079 19.940 ATOM 1853 CD1 LEU A 242 115.390 53.933 18.970 1.00 37.59 CD2 LEU A 242 **ATOM** 1854 113.748 54.825 20.654 1.00 37.84 PHE A 243 1855 55.209 MOTA N 118.785 22.666 1.00 36.35 MOTA 1856 PHE A 243 119.749 55.879 23.534 1.00 36.90 CA PHE A 243 MOTA 1857 121.143 55.806 C 22.930 1.00 37.73 PHE A 243 **ATOM** 1858 121.890 56.781 22.958 1.00 38.96 0 **ATOM** 1859 119.769 55.262 CB PHE A 243 24.938 1.00 35.66 55.683 **MOTA** 1860 25.803 CG PHE A 243 118.619 1.00 34.47 1861 118.325 57.032 25.980 1.00 34.10 CD1 PHE A 243 MOTA CD2 PHE A 243 MOTA 1862 117.850 54.739 26.474 1.00 33.97 CE1 PHE A 243 ATOM 1863 117.280 57.432 26.818 1.00 33.89 1864 CE2 PHE A 243 116.802 55.130 1.00 33.03 **ATOM** 27.317 1865 CZ PHE A 243 116.517 56.474 27.489 1.00 32.69 ATOM 1866 121.477 **GLU A 244** 54.650 22.370 1.00 38.32 MOTA N **GLU A 244** ATOM 1867 CA 122.779 54.431 21.763 1.00 39.15 122.974 **GLU A 244** 55.304 20.531 1.00 37.76 **ATOM** 1868 C 124.103 55.550 **GLU A 244** 20.103 1.00 37.31 MOTA 1869 MOTA **GLU A 244** 122.932 52,959 21.380 1.00 43.06 1870 CB **GLU A 244** 124.297 MOTA 52.595 20.823 1871 CG 1.00 48.69 **GLU A 244** 51.141 **ATOM** 1872 CD124.375 20.407 1.00 53.07 50.272 1873 124.097 ATOM OE1 GLU A 244 21.266 1.00 55.43 124.713 OE2 GLU A 244 50.866 19.228 1.00 54.65 MOTA 1874 55.767 1.00 36.29 MOTA 1875 ALA A 245 19.951 121.873 N ALA A 245 18.770 1.00 35.13 MOTA 1876 CA 121.964 56.619 1877 58.061 19.224 1.00 33.93 ALA A 245 122.120 ATOM C 1878 58.935 18.437 1.00 34.56 ALA A 245 122.455 MOTA 120.723 56.463 MOTA ALA A 245 17.897 1.00 35.38 1879 1.00 32.96 121.873 58.301 **GLY A 246** 20.506 MOTA 1880 N **GLY A 246** 59.640 1.00 32.17 21.039 MOTA 1881 CA 122.022 60.256 21.720 1.00 31.72 **GLY A 246** 120.812 MOTA 1882 C 22.265 1.00 32.01 ATOM 1883 120.919 61.358 GLY A 246 MOTA 1884 119.673 59.570 21.710 1.00 30.21 ALA A 247 N 1.00 29.83 118.465 22.328 1885 ALA A 247 60.123 CA MOTA 1.00 29.13 ALA A 247 60.676 23.718 1886 C 118.739 ATOM 24.529 1.00 29.36 119.392 60.038 ALA A 247 ATOM 1887 117.365 1.00 29.66 59.064 22.398 MOTA 1888 CB ALA A 247 118.231 61.874 23.976 1.00 29.42 N **ASP A 248** MOTA 1889

### 62.541 MOTA 1890 CA ASP A 248 118.401 25.262 1.00 29.22 C **ASP A 248** 62.141 26.218 MOTA 1891 C 117.283 1.00 28.93 C 1892 MOTA 0 **ASP A 248** 117.334 62.412 27.416 1.00 28.35 0 MOTA 1893 **ASP A 248** 1.00 31.00 CB 118.400 64.049 25.032 C **ASP A 248** MOTA 1894 CG 119.557 64.495 C 24.153 1.00 33.31 MOTA 1895 120.674 64.660 OD1 ASP A 248 1.00 34.76 24.689 0 OD2 ASP A 248 1896 **ATOM** 119.359 64.648 22.922 1.00 33.85 0 **ATOM** 1897 ALA A 249 61.487 25.666 1.00 28.79 N 116.268 N 1898 ALA A 249 115.127 61.020 26.439 ATOM CA 1.00 28.18 C 114.279 60.097 1899 ALA A 249 25.568 C ATOM C 1.00 28.88 ALA A 249 114.314 **ATOM** 1900 0 60.181 24.341 1.00 28.79 0 ALA A 249 1901 114.298 62.209 26.903 1.00 27.25 C MOTA CB **ATOM** 1902 ILE A 250 113.543 59.189 26.192 1.00 28.48 N N 1903 ILE A 250 112.669 58.324 25.421 1.00 29.55 C MOTA CA ILE A 250 111.260 25.916 1.00 29.65 MOTA 1904 C 58.577 C ILE A 250 1905 111.056 58.932 27.085 1.00 29.05 0 0 MOTA ILE A 250 MOTA 1906 CB 113.000 56.813 25.587 1.00 30.94 C CG1 ILE A 250 1907 113.023 56.441 27.070 1.00 32.35 C MOTA CG2 ILE A 250 114.303 1.00 30.20 C MOTA 1908 56.479 24.876 CD1 ILE A 250 C 1909 113.296 54.968 27.328 1.00 34.16 MOTA VAL A 251 110.289 58.414 25.024 1.00 29.98 MOTA 1910 N N VAL A 251 108.894 58.615 25.385 C MOTA CA 1911 1.00 30.11 VAL A 251 1912 108.119 1.00 31.15 C MOTA C 57.310 25.254 VAL A 251 56.777 **ATOM** 0 1913 0 107.939 24.157 1.00 31.33 1914 VAL A 251 108.222 59.698 24.499 C **ATOM** CB 1.00 29.28 CG1 VAL A 251 C 59.837 1.00 28.74 1915 106.749 24.869 MOTA CG2 VAL A 251 108.924 61.032 1.00 28.73 C **ATOM** 1916 24.689 1917 ILE A 252 56.791 26.394 N ATOM 107.686 1.00 32.25 N 1.00 34.36 **ATOM** 1918 CA ILE A 252 106.907 55.567 26.445 ILE A 252 105.488 56.052 26.169 1.00 35.92 C **ATOM** 1919 C 104.796 27.055 0 MOTA 1920 0 ILE A 252 56.541 1.00 35.07 1.00 34.93 27.838 1921 54.933 C ILE A 252 107.030 MOTA CB C CG1 ILE A 252 108.516 28.119 1.00 34.36 MOTA 1922 54.658 CG2 ILE A 252 106.190 C MOTA 53.662 27.918 1.00 34.35 1923 C CD1 ILE A 252 1.00 35.76 54.276 MOTA 1924 108.830 29.536 105.084 N 1925 55.928 24.911 1.00 37.82 ASP A 253 MOTA N C ASP A 253 1.00 38.30 MOTA 1926 ÇA 103.793 56.405 24.439 102.759 C 1927 **ASP A 253** 55.298 24.528 1.00 36.33 ATOM C 102.900 0 54.256 23.897 ASP A 253 1.00 35.52 MOTA 1928 0 56.886 1.00 42.68 C ASP A 253 103.969 MOTA 1929 22.987 CB C 103.027 58.024 1930 ASP A 253 22.608 1.00 47.26 MOTA CG 0 1.00 49.54 1931 103.007 59.059 23.324 OD1 ASP A 253 ATOM 0 102.326 57.895 1.00 49.50 1932 21.574 ATOM OD2 ASP A 253 THR A 254 101.720 55.535 25.319 1.00 35.67 N 1933 N ATOM C 25.507 1.00 35.99 1934 100.654 54.553 ATOM CA THR A 254 C 1935 THR A 254 99.267 55.208 25.434 1.00 33.80 ATOM C 0 99.115 56.398 25.716 1.00 33.46 THR A 254 ATOM 1936 0

### 100.769 53.870 26.894 CB THR A 254 MOTA 1937 1.00 37.36 C OG1 THR A 254 1.00 40.12 MOTA 1938 102.127 53.489 27.131 0 CG2 THR A 254 **ATOM** 1939 1.00 39.58 C 99.898 52.617 26.943 **ALA A 255** 98.257 N 25.076 1.00 32.56 N **ATOM** 1940 54.424 ALA A 255 96.901 1.00 31.89 C 1941 54.943 ATOM CA 25.000 ALA A 255 96.381 55.122 1.00 31.60 C 1942 C 26.419 MOTA ALA A 255 95.756 56.131 1.00 31.27 0 MOTA 1943 0 26.743 ALA A 255 96.021 53.983 C 24.239 1.00 31.75 1944 CB ATOM 1.00 31.05 HIS A 256 N **MOTA** 1945 N 96.646 54.133 27.266 96.211 28.652 1.00 30.66 C HIS A 256 54.183 MOTA 1946 CA 53.961 C HIS A 256 97.423 29.541 1947 C 1.00 31.22 MOTA HIS A 256 29.784 1.00 32.42 0 52.826 **ATOM** 0 97.811 1948 1.00 28.92 HIS A 256 95.183 53.092 C 1949 28.932 **ATOM** CB 1.00 28.92 C HIS A 256 94.514 53.230 1950 30.263 CG MOTA ND1 HIS A 256 93.767 52.223 30.832 N **ATOM** 1951 1.00 29.33 1.00 28.44 C CD2 HIS A 256 94.461 54.270 1952 31.128 MOTA C **CE1 HIS A 256** 93.284 52.636 31.991 1.00 29.11 1953 MOTA NE2 HIS A 256 93.690 32.194 1.00 29.27 N 1954 **ATOM** 53.875 1.00 32.56 1955 **GLY A 257** 98.017 55.043 30.026 И N ATOM 54.927 C **GLY A 257** 1.00 33.03 MOTA 99.196 30.871 1956 CA C 1957 **GLY A 257** 98.907 54.494 32.297 1.00 33.73 ATOM C 33.073 **GLY A 257** 99.825 54.233 0 1.00 34.44 MOTA 1958 0 32.655 1.00 33.24 1959 HIS A 258 97.633 54.407 И **ATOM** N $\mathsf{C}$ 97.277 54.009 34.005 1.00 32.25 HIS A 258 MOTA 1960 CA C 1961 HIS A 258 97.016 52.500 33.995 1.00 33.33 MOTA C 1.00 34.50 0 HIS A 258 51.991 1962 96.237 34.796 MOTA 1.00 30.67 C 1963 HIS A 258 96.025 54.768 34.447 **MOTA** CB C HIS A 258 95.892 54.923 1964 35.932 1.00 28.53 ATOM CG ND1 HIS A 258 N 96.576 36.831 1.00 28.44 **ATOM** 54.135 1965 1.00 27.04 C CD2 HIS A 258 95.100 55.735 36.672 **ATOM** 1966 **CE1 HIS A 258** 38.059 MOTA 1967 96.212 54.455 1.00 27.92 55.422 N 37.991 1.00 27.43 NE2 HIS A 258 95.316 MOTA 1968 51.790 1.00 33.64 Ν 97.664 33.076 ATOM SER A 259 1969 N C SER A 259 97.502 32.974 1.00 35.31 50.340 1970 CA ATOM C 33.765 1.00 35.20 98.572 49.598 SER A 259 ATOM 1971 99.745 SER A 259 33.751 1.00 35.19 1972 49.975 MOTA C 97.593 49.883 1.00 36.71 SER A 259 31.521 **ATOM** 1973 CB 0 30.734 1.00 42.41 MOTA 1974 SER A 259 96.600 50.498 OG N 48.525 34.434 1.00 35.34 ALA A 260 1975 98.160 MOTA N C 35.228 1.00 34.81 MOTA ALA A 260 99.075 47.718 1976 CA C 100.264 1.00 34.54 ALA A 260 47.288 34.377 1977 ATOM C 0 47.394 34.797 101.415 1.00 34.57 ATOM ALA A 260 1978 C 98.345 1.00 34.40 46.500 35.769 **ATOM** ALA A 260 1979 CB1.00 34.40 N 33.174 1980 GLY A 261 99.971 46.808 MOTA N C GLY A 261 1.00 33.69 46.368 32.271 101.013 MOTA 1981 CA 1.00 33.68 C 102.003 31.920 MOTA GLY A 261 47.458 1982 C 31.923 1.00 35.39 103.211 47.220 1983 0 GLY A 261 ATOM

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### **ATOM** 1984 VAL A 262 N 101.505 48.652 31.612 1.00 31.87 N **ATOM** VAL A 262 1985 CA 102.372 49.769 31.260 1.00 30.22 C 1986 VAL A 262 MOTA C 103.257 50.106 32.453 1.00 30.85 C **ATOM** VAL A 262 1987 0 104.459 50.296 32.310 1.00 29.76 0 VAL A 262 MOTA 1988 101.545 CB 51.037 30.851 1.00 29.14 C **ATOM** 1989 CG1 VAL A 262 102.469 52.200 30.517 1.00 25.83 C **ATOM** 1990 CG2 VAL A 262 100.668 29.653 1.00 28.66 50.723 C **ATOM** 1991 **LEU A 263** N 102.657 50.170 1.00 32.42 33.635 N 1992 **LEU A 263** ATOM CA 103.404 50.504 34.842 1.00 35.12 C ATOM 1993 LEU A 263 C 104.479 49.449 35.109 C 1.00 37.28 ATOM 1994 **LEU A 263** 0 105.579 49.747 35.580 1.00 37.10 0 LEU A 263 102.445 50.611 MOTA 1995 CB 36.035 1.00 33.85 C MOTA 1996 101.288 CG LEU A 263 35.876 51.607 C 1.00 32.67 CD1 LEU A 263 MOTA 1997 100.436 51.594 37.126 1.00 31.79 C MOTA 1998 CD2 LEU A 263 101.827 53.004 35.615 C 1.00 31.44 104.148 ARG A 264 MOTA 1999 N 48.208 34.792 1.00 39.38 N MOTA 2000 ARG A 264 CA 47.107 105.067 34.981 1.00 41.52 C C 47.304 MOTA 2001 ARG A 264 106.272 34.064 C 1.00 40.90 ARG A 264 MOTA 2002 107.410 0 34.517 47.312 1.00 40.25 0 ARG A 264 MOTA 2003 CB 104.329 45.804 34.674 1.00 45.25 C MOTA ARG A 264 2004 105.121 CG 34.813 1.00 50.16 44.513 C MOTA 2005 CDARG A 264 104.144 43.343 34.780 C 1.00 53.35 ARG A 264 103.248 43.420 MOTA 2006 NE 1.00 56.71 33.626 N **MOTA** 2007 ARG A 264 CZ102.152 42.681 33.479 1.00 58.23 C NH1 ARG A 264 101.810 ATOM 2008 41.805 N 34.416 1.00 60.06 NH2 ARG A 264 ATOM 2009 101.390 42.821 32.402 1.00 58.84 N LYS A 265 106.010 47.486 ATOM 2010 N 32.776 1.00 40.84 N LYS A 265 ATOM 2011 CA 107.068 47.684 31.795 1.00 41.03 C MOTA 2012 LYS A 265 C 107.915 48.928 1.00 41.03 C 32.068 LYS A 265 48.909 31.885 MOTA 2013 109.130 1.00 42.30 0 MOTA LYS A 265 2014 CB 106.461 47.762 C 30.390 1.00 41.68 2015 ATOM CG LYS A 265 107.441 48.146 29.293 1.00 43.10 C MOTA 2016 LYS A 265 108.624 C CD47.198 29.235 1.00 45.73 108.175 MOTA LYS A 265 45.761 C 2017 CE 29.019 1.00 47.46 **ATOM** LYS A 265 N 2018 NZ107.382 45.593 27.771 1.00 48.86 N ATOM 2019 107.278 32.504 N ILE A 266 50.009 1.00 40.38 ILE A 266 1.00 39.51 C MOTA 2020 CA 107.999 51.244 32.783 C ATOM 2021 108.989 51.061 C ILE A 266 33.920 1.00 40.45 **ATOM** 2022 ILE A 266 110.124 51.549 33.856 Q 1.00 40.08 0 ILE A 266 107.028 C ATOM 2023 52.396 33.134 CB 1.00 38.28 2024 106.249 52.809 31.887 1.00 36.55 C CG1 ILE A 266 ATOM 1.00 37.78 C CG2 ILE A 266 53.572 ATOM 2025 107.795 33.704

105.333

108.553

109.404

110.584

111.704

53.963

50.359

50.105

49.260

49.453

32.111

34.962

36.117

35.672

36.141

1.00 36.83

1.00 41.07

1.00 41.67

1.00 42.17

1.00 42.01

ΑT	OM 2	2031	CB	ALA	A	267	108.621	49.386	37.200	1.00	41.62	C
AT	OM 2	2032	N	GLU	Α	268	110.330	48.330	34.757	1.00	42.65	N
			CA	GLU	Α	268	111.386	47.470	34.246			C
				GLU			112.334	48.294	33.383		44.45	C
				GLU			113.548	48.064	33.382	1.00		0
		2036		GLU			110.790	46.321	33.429			Ċ
				GLU			109.741	45.530	34.195		50.26	Č
		2037				268	109.164	44.377	33.394	1.00		C
				GLU			103.776	44.587	32.222		53.91	0
						268	109.080	43.260	33.947		54.82	0
			N				111.780	49.256	32.649		43.95	N
				ILE								C
		2042	CA	ILE			112.591	50.117	31.799		43.12	C
		2043	C	ILE			113.450	51.012	32.688		42.99	
			0	ILE			114.627	51.236	32.402		42.66	0
		2045	CB	ILE			111.716	51.001	30.881		42.99	C
		2046	CG1	ILE			110.944	50.122	29.896		42.46	C
		2047	CG2	ILE			112.590	51.995	30.123		42.46	C
ΑŢ	COM 2	2048	CD1	ILE			110.036	50.898	28.968		41.32	C
ΓA	COM 2	2049	N	ARG			112.859	51.519	33.768		42.35	N
ΓA	COM 2	2050	CA	ARG	A	270	113.592	52.372	34.696		42.43	C
ΓA	COM 2	2051	C	ARG	A	270	114.667	51.580	35.447		43.30	C
ΑT	OM 2	2052	0	ARG	A	270	115.802	52.033	35.598	1.00	42.17	0
ΓA	COM 2	2053	CB	ARG	A	270	112.630	53.026	35.689	1.00	40.28	C
ΑT	OM 2	2054	CG	ARG	A	270	113.335	53.693	36.857	1.00	40.08	C
AT	COM 2	2055	CD	ARG	A	270	114.401	54.661	36.381	1.00	39.52	C
AT	OM 2	2056	NE	ARG	A	270	113.843	55.890	35.835	1.00	39.63	N
ΑT	OM 2	2057	CZ	ARG	Α	270	114.568	56.833	35.246	1.00	40.17	C
ΓA	OM 2	2058	NH1	ARG	A	270	115.879	56.682	35.127	1.00	39.75	N
ΓA	OM 2	2059	NH2	ARG	A	270	113.991	57.941	34.803	1.00	41.04	N
ΓA	OM 2	2060	N	ALA	Α	271	114.297	50.393	35.917	1.00	44.99	N
		2061	CA	ALA	Α	271	115.224	49.538	36.638	1.00	46.51	C
		2062	С	ALA	Α	271	116.445	49.281	35.766	1.00	47.60	C
		2063	0	ALA	Α	271	117.550	49.086	36.272	1.00	49.90	0
		2064	CB	ALA			114.555	48.222	36.998	1.00	46.57	C
		2065	N	HIS	A	272	116.254	49.283	34.454	1.00	47.42	N
		2066	CA			272	117.368	49.048	33.559	1.00	47.00	С
		2067	C	HIS			118.096	50.347	33.204	1.00	46.88	С
		2068	0	HIS			119.308	50.339	33.009	1.00	47.71	0
		2069	CB	HIS			116.889	48.341	32.295	1.00	47.98	C
		2070	CG	HIS		272	117.999	47.783	31.461	1.00	50.44	C
		2071	ND1	HIS		272	117.779	47.130	30.269		51.34	N
		2072	CD2	HIS			119.341	47.762	31.658	1.00		С
		2072	CE1			272	118.935	46.730	29.767	1.00		C
		2073	NE2			272	119.899	47.101	30.591		51.04	N
		207 <del>4</del> 2075	NEZ			273	117.370		33.124		46.29	N
		2075	CA			273	117.991		32.806		45.64	C
			CA			273	117.810		33.954		46.32	C
A'	rom :	2077	C	rne	A	413	111.010	JJ.1JL	JJ • JJ <del>•</del>	1.00	10.00	•

ATOM	2078	0	PHE	A	273	117.128	54.770	33.797	1.00	47.02	0
ATOM	2079	CB	PHE	Α	273	117.386	53.378	31.541	1.00	43.67	C
ATOM	2080	CG	PHE	A	273	117.491	52.519	30.313	1.00	42.54	C
ATOM	2081	CD1	PHE	Α	273	116.589	51.489	30.088	1.00	42.71	C
ATOM	2082	CD2	PHE	Α	273	118.485	52.755	29.370	1.00	42.26	C
ATOM	2083	CE1	PHE	A	273	116.673	50.709	28.933	1.00	43.23	C
ATOM	2084	CE2	PHE	Α	273	118.580	51.981	28.217	1.00	41.40	C
ATOM	2085	CZ	PHE	Α	273	117.673	50.959	27.996	1.00	42.59	C
ATOM	2086	N	PRO	Α	274	118.431	53.483	35.116	1.00	45.94	N
ATOM	2087	CA	PRO	Α	274	118.355	54.329	36.313	1.00	45.40	C
ATOM	2088	C	PRO	A	274	118.763	55.799	36.194	1.00	45.24	C
ATOM	2089	0	PRO	Α	274	118.347	56.614	37.011	1.00	45.24	0
ATOM	2090	CB	PRO	Α	274	119.229	53.571	37.308	1.00	44.99	C
ATOM	2091	CG	PRO	A	274	120.244	52.905	36.395	1.00	44.50	C
MOTA	2092	CD	PRO	A	274	119.283	52.317	35.400	1.00	45.59	C
MOTA	2093	N	ASN	A	275	119.566	56.145	35.195	1.00	45.09	N
MOTA	2094	CA	ASN	A	275	120.015	57.527	35.042	1.00	45.14	C
MOTA	2095	C	ASN	A	275	119.229	58.295	33.977	1.00	44.49	C
MOTA	2096	0	ASN	Α	275	118.967	59.487	34.126	1.00	45.55	0
MOTA	2097	CB	ASN	A	275	121.497	57.563	34.664	1.00	47.44	С
ATOM	2098	CG	ASN	A	275	122.370	56.762	35.614	1.00	49.31	C
MOTA	2099	OD1	ASN	A	275	122.481	57.076	36.795	1.00	49.47	0
MOTA	2100	ND2	ASN	A	275	123.000	55.713	35.090	1.00	51.15	N
MOTA	2101	N	ARG	A	276	118.869	57.600	32.903	1.00	41.92	N
MOTA	2102	CA	ARG	Α	276	118.143	58.183	31.782	1.00	39.15	C
MOTA	2103	С	ARG	A	276	116.868	58.946	32.148	1.00	37.69	С
MOTA	2104	0	ARG	A	276	116.276	58.729	33.206	1.00	37.58	0
ATOM	2105	CB	ARG	A	276	117.793	57.080	30.791	1.00	39.18	C
ATOM	2106	CG	ARG	A	276	118.985	56.271	30.302	1.00	37.60	C
MOTA	2107	CD	ARG	Α	276	119.955	57.089	29.463	1.00	36.19	C
MOTA	2108	NE	ARG	Α	276	121.019	56.235	28.938	1.00	35.76	N
MOTA	2109	CZ	ARG	Α	276	121.913	56.609	28.031	1.00	35.25	C
ATOM	2110	NH1	ARG	A	276	121.887	57.836	27.531	1.00	36.02	N
MOTA	2111	NH2	ARG	Α	276	122.831	55.750	27.617	1.00	36.01	N
MOTA	2112	N	THR	A	277	116.461	59.846	31.254	1.00	35.95	И
ATOM	2113	CA	THR	Α	277	115.251	60.654	31.429	1.00	33.86	C
MOTA	2114	С	THR	Α	277	114.086	59.917	30.790	1.00	32.47	C
MOTA	2115	0	THR	A	277	114.108	59.638	29.587	1.00	31.01	0
ATOM	2116	CB	THR	Α	277	115.369	62.023	30.728	1.00	33.80	C
MOTA	2117	OG1	THR	Α	277	116.362	62.822	31.381	1.00	34.95	0
MOTA	2118	CG2	THR	A	277	114.039	62.746	30.753	1.00	33.78	C
MOTA	2119	N	LEU	A	278	113.073	59.602	31.588	1.00	30.23	N
ATOM	2120	CA	LEU	A	278	111.915	58.889	31.070	1.00	29.28	C
MOTA	2121	C	LEU	A	278	110.663	59.754	31.048	1.00	28.27	C
MOTA	2122	0	LEU	Α	278	110.329	60.404	32.032	1.00	27.31	0
ATOM	2123	СВ	LEU	A	278	111.649	57.628	31.893	1.00	28.56	C
MOTA	2124	CG	LEU	A	278	112.782	56.600	31.915	1.00	29.99	C

ATOM	2125	CD1	LEU	A	278	112.339	55.362	32.692	1.00	29.20	C
MOTA	2126	CD2	LEU	A	278	113.159	56.217	30.486	1.00	30.32	C
MOTA	2127	N	ILE	Α	279	109.993	59.766	29.900	1.00	26.83	N
MOTA	2128	CA	ILE	A	279	108.759	60.517	29.713	1.00	25.75	C
MOTA	2129	C	ILE	A	279	107.756	59.403	29.451	1.00	26.77	C
ATOM	2130	0	ILE	A	279	108.035	58.499	28.649	1.00	25.38	0
ATOM	2131	CB	ILE	A	279	108.847	61.448	28.485	1.00	23.44	C
MOTA	2132	CG1	ILE	A	279	110.078	62.345	28.598	1.00	23.15	C
MOTA	2133	CG2	ILE	A	279	107.604	62.310	28.405	1.00	24.07	C
MOTA	2134	CD1	ILE	A	279	110.295	63.233	27.412	1.00	23.91	C
MOTA	2135	N	ALA			106.596	59.449	30.100	1.00	27.48	N
MOTA	2136	CA	ALA			105.657	58.354	29.914	1.00		C
MOTA	2137	C	ALA			104.246	58.667	29.469	1.00	32.08	C
MOTA	2138	0			280	103.726	59.745	29.725	1.00	31.46	0
ATOM	2139	CB	ALA			105.608	57.495	31.182	1.00	30.65	C
ATOM	2140	N	GLY			103.665	57.644	28.829	1.00	35.24	N
ATOM	2141	CA	GLY			102.314	57.614	28.273	1.00	35.20	C
MOTA	2142	C	GLY			101.303	58.614	28.763	1.00	35.43	C
MOTA	2143	0	GLY			101.623	59.501	29.552	1.00	37.45	0
MOTA	2144	N	ASN			100.057	58.467	28.325	1.00	32.74	N
ATOM	2145	CA	ASN			99.063	59.443	28.729	1.00	29.41	С
MOTA	2146	C	ASN			98.088	59.056	29.815	1.00	27.37	C
MOTA	2147	0	ASN			97.590	57.933	29.882		26.60	0
MOTA	2148	CB	ASN			98.318	59.939	27.491	1.00	30.24	C
ATOM	2149	CG	ASN			99.200	60.807	26.595		31.84	C
ATOM	2150	OD1				100.405	60.562	26.467		32.15	0
ATOM	2151	ND2	ASN			98.600	61.809	25.949	1.00		N
MOTA	2152	N			283	97.845	60.019	30.690		25.51	N
ATOM	2153	CA			283	96.907	59.873	31.790		24.37	C
ATOM	2154	C			283	96.136	61.185	31.823		23.61	C
MOTA	2155	0			283	96.532	62.160	31.175		23.14	0
MOTA	2156	СВ			283	97.636	59.631	33.150		23.62	C
ATOM	2157	CG1			283	98.677	60.727	33.421		22.08	C
MOTA	2158	CG2			283	98.279	58.256	33.153		23.25	C
MOTA	2159	CD1			283	98.119	62.094	33.881		19.86	C
MOTA	2160	N			284	95.039	61.216	32.561		22.29	N
MOTA	2161	CA			284	94.254	62.426	32.639		23.16	C
ATOM	2162	C			284	93.744	62.633	34.059		24.19	C
ATOM	2163	0			284	92.839	63.434	34.300		25.24	0
MOTA	2164	CB			284	93.102	62.347	31.659		22.62	C
MOTA	2165	N			285	94.341	61.914	35.004		25.00	N
ATOM	2166	CA			285	93.936	62.015	36.403		25.00	C
ATOM	2167	C			285	95.149	62.180	37.308		24.68	C
ATOM	2168	0			285	96.282	61.956	36.889		22.17	0
MOTA	2169				285		60.759	36.851		24.60	C
MOTA	2170		THR			94.056	59.636	36.784		27.11	0
MOTA	2171	CG2	THR	A	285	91.975	60.501	35.953	1.00	23.14	C

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ATOM	2219	N	TYR	A	293	104.661	58.408	38.083	1.00 27.07	N
MOTA	2220	CA	TYR	A	293	105.809	58.947	38.804	1.00 28.43	C
MOTA	2221	C	TYR	A	293	106.353	57.853	39.710	1.00 29.43	C
MOTA	2222	0	TYR	Α	293	107.555	57.601	39.738	1.00 29.87	0
MOTA	2223	CB	TYR	Α	293	105.398	60.155	39.629	1.00 28.36	C
MOTA	2224	CG	TYR	A	293	105.192	61.411	38.812	1.00 29.78	C
MOTA	2225	CD1	TYR	A	293	104.175	62.310	39.134	1.00 29.30	C
MOTA	2226	CD2	TYR	A	293	106.038	61.725	37.744	1.00 29.08	C
MOTA	2227	CE1	TYR	A	293	104.000	63.485	38.417	1.00 28.70	C
ATOM	2228	CE2	TYR	A	293	105.868	62.905	37.020	1.00 28.90	C
MOTA	2229	CZ	TYR	A	293	104.844	63.776	37.367	1.00 28.75	C
MOTA	2230	OH	TYR			104.644	64.936	36.666	1.00 29.75	0
MOTA	2231	N	ASP			105.456	57.191	40.435	1.00 30.73	N
MOTA	2232	CA			294	105.846	56.104	41.325	1.00 32.03	C
ATOM	2233	C			294	106.533	54.961	40.573	1.00 33.10	C
MOTA	2234	0			294	107.287	54.185	41.167	1.00 34.97	0
ATOM	2235	CB	ASP		294	104.623	55.570	42.067	1.00 31.61	С
ATOM	2236	CG	ASP			104.081	56.554	43.080	1.00 32.26	C
ATOM	2237	OD1				103.034	56.266	43.700	1.00 33.58	0
ATOM	2238	OD2	ASP			104.704	57.615	43.267	1.00 32.44	0
ATOM	2239	N	ALA			106.275	54.854	39.271	1.00 32.83	N
ATOM	2240	CA	ALA			106.879	53.802	38.455	1.00 31.74	C
ATOM	2241	C	ALA			108.264	54.224	37.992	1.00 31.68	C
ATOM	2242	0	ALA			108.985	53.436	37.380	1.00 32.56	0
ATOM	2243	CB	ALA			105.997	53.482	37.251	1.00 30.93	C
MOTA	2244	N	GLY			108.630	55.470	38.269	1.00 30.95	N
ATOM	2245	CA	GLY			109.948	55.933	37.882 36.717	1.00 31.13 1.00 31.66	C
ATOM	2246	С 0	GLY			110.045 111.149	56.900 57.204	36.265	1.00 31.00	0
ATOM ATOM	2247 2248	N	VAL			108.919	57.394	36.219	1.00 31.51	N
ATOM	2249	CA	VAL		297	108.978	58.328	35.101	1.00 31.83	C
ATOM	2250	C	VAL			109.416	59.716	35.586	1.00 31.03	C
ATOM	2251	0	VAL			109.171	60.092	36.730	1.00 29.21	0
ATOM	2252	СВ			297	107.606	58.445	34.382	1.00 32.91	C
ATOM	2253	CG1				106.596	59.096	35.290	1.00 35.68	С
ATOM	2254	CG2	VAL			107.747	59.266	33.125	1.00 35.70	С
ATOM	2255	N	ASP		298	110.074	60.463	34.707	1.00 28.22	N
MOTA	2256	CA	ASP	Α	298	110.545	61.801	35.022	1.00 27.00	C
ATOM	2257	С	ASP	A	298	109.528	62.852	34.607	1.00 27.20	C
ATOM	2258	0	ASP	Α	298	109.281	63.813	35.332	1.00 26.86	0
MOTA	2259	CB	ASP	A	298	111.870	62.071	34.309	1.00 28.58	C
MOTA	2260	CG	ASP	Α	298	112.974	61.149	34.776	1.00 30.31	C
ATOM	2261	OD1	ASP	A	298	113.221	61.114	35.994	1.00 34.37	0
ATOM	2262	OD2	ASP	A	298	113.600	60.462	33.943	1.00 31.52	0
ATOM	2263	N	VAL	A	299	108.942	62.664	33.430	1.00 26.94	N
ATOM	2264	CA	VAL	A	299	107.955	63.596	32.898	1.00 25.40	C
ATOM	2265	С	VAL	A	299	106.704	62.825	32.462	1.00 24.71	C

### MOTA 2266 0 VAL A 299 106.792 61.831 31.736 1.00 24.19 0 2267 VAL A 299 108.526 64.371 31.667 1.00 25.18 ATOM CB C **ATOM** 2268 CG1 VAL A 299 31.255 107.573 65.493 1.00 23.30 C CG2 VAL A 299 31.985 1.00 24.81 C MOTA 2269 109.908 64.920 **VAL A 300** 2270 105.544 63.287 1.00 23.99 N ATOM N 32.916 **VAL A 300** 2271 CA 104.270 62.661 32.576 1.00 23.71 C MOTA MOTA 2272 **VAL A 300** 31.555 C C 103.499 63.493 1.00 24.32 **VAL A 300** MOTA 2273 103.338 64.707 31.730 1.00 24.20 MOTA 2274 CB **VAL A 300** 103.370 33.825 1.00 23.42 C 62.486 CG1 VAL A 300 62.116 1.00 23.29 C **ATOM** 2275 101.967 33.407 2276 CG2 VAL A 300 103.923 61.391 34.722 1.00 23.23 C ATOM LYS A 301 62.854 2277 103.033 1.00 24.40 ATOM N 30.484 N 2278 LYS A 301 63.588 **ATOM** 1.00 24.04 C CA 102.258 29.503 LYS A 301 100.782 1.00 22.78 MOTA 2279 C 63.356 29.753 C LYS A 301 2280 100.322 62.229 29.960 1.00 21.52 0 ATOM 0 LYS A 301 **ATOM** 2281 CB 102.686 63.259 28.065 1.00 24.29 C 2282 LYS A 301 61.818 27.709 C MOTA ÇG 102.809 1.00 29.51 C LYS A 301 103.757 61.674 26.489 1.00 32.99 ATOM 2283 CD LYS A 301 103.228 C 2284 62.284 25.187 1.00 30.67 MOTA CE LYS A 301 2285 102.178 61.439 24.564 1.00 30.01 N MOTA NZ**VAL A 302** 100.060 64.470 1.00 22.18 29.782 N 2286 N MOTA C **ATOM** 2287 VAL A 302 98.641 64.485 30.061 1.00 22.10 CA **VAL A 302** 64.628 28.807 1.00 22.59 C **ATOM** C 97.793 2288 MOTA **VAL A 302** 98.082 65.457 27.938 1.00 22.52 0 2289 0 C VAL A 302 CB 98.306 65.652 31.032 1.00 21.25 ATOM 2290 CG1 VAL A 302 C **ATOM** 96.815 65.677 31.335 1.00 20.58 2291 CG2 VAL A 302 1.00 20.42 C 2292 99.103 65.504 32.312 MOTA **GLY A 303** 1.00 22.97 96.745 63.812 28.722 N ATOM 2293 N **GLY A 303** 63.892 27.582 1.00 22.25 C 95.854 MOTA 2294 CA C **GLY A 303** 62.618 27.153 2295 C 95.160 1.00 22.34 MOTA 95.784 61.750 26.566 ATOM 2296 1.00 23.71 **GLY A 303** 0 N 93.873 62.493 27.453 1.00 22.67 ILE A 304 MOTA 2297 N 93.107 61.329 C 27.011 1.00 24.68 ILE A 304 MOTA 2298 CA 91.883 61.799 C MOTA ILE A 304 26.202 1.00 26.38 2299 C 90.881 1.00 26.11 0 62.237 26.770 MOTA ILE A 304 2300 0 1.00 23.24 C 92.625 60.446 28.195 ILE A 304 MOTA 2301 CBC 28.906 1.00 22.41 2302 93.818 59.799 MOTA CG1 ILE A 304 C 59.346 27.680 1.00 21.36 MOTA 2303 CG2 ILE A 304 91.711 C 28.113 94.503 58.718 1.00 22.43 MOTA 2304 CD1 ILE A 304 N GLY A 305 91.996 61.747 24.872 1.00 28.56 2305 N MOTA C 90.895 62.138 1.00 30.63 2306 24.011 ATOM CA GLY A 305 23.356 GLY A 305 90.843 63.516 1.00 33.34 MOTA 2307 C 0 63.666 1.00 34.88 90.112 22.370 **ATOM** 2308 0 GLY A 305 N 91.588 64.534 23.836 1.00 33.14 2309 PRO A 306 ATOM N 23.226 91.541 1.00 33.01 65.875 **ATOM** 2310 CA PRO A 306 92.126 66.054 21.814 1.00 33.06 C PRO A 306 ATOM 2311 91.928 67.104 21.191 1.00 33.29 2312 0 PRO A 306 ATOM

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### ATOM 2313 CB PRO A 306 92.275 66.728 24.256 1.00 30.93 C MOTA 2314 PRO A 306 93.353 65.772 CG 24.704 1.00 30.96 C MOTA 2315 PRO A 306 64.547 CD 92.504 24.992 C 1.00 31.97 **ATOM** 2316 **GLY A 307** 92.833 N 21.313 65.044 1.00 33.26 N **ATOM** 2317 CA **GLY A 307** 93.446 65.148 19.994 1.00 33.13 C **ATOM** 2318 **GLY A 307** 92.525 65.502 C 18.837 1.00 33.48 C MOTA 2319 0 **GLY A 307** 91.414 64.982 18.739 1.00 33.64 0 **ATOM** 2320 SER A 308 92.992 66.382 17.950 N 1.00 32.89 N MOTA 2321 **SER A 308** 92.209 66.802 16.783 CA C 1.00 33.08 **ATOM** 2322 **SER A 308** C 91.745 65.606 15.950 C 1.00 32.80 **ATOM** 2323 **SER A 308** 90.623 0 65.577 15.452 0 1.00 33.12 2324 **SER A 308** 93.038 15.898 MOTA CB 67.743 1.00 32.26 C 2325 **SER A 308** 94.229 ATOM OG 1.00 31.53 67.114 15.448 0 **ATOM** 2326 ILE A 309 92.625 N 64.625 15.809 1.00 33.27 N **MOTA** 2327 ILE A 309 63.416 CA 92.346 C 15.045 1.00 34.38 **ATOM** 2328 C ILE A 309 91.953 62.229 15.940 1.00 34.76 C 2329 ILE A 309 91.843 61.095 ATOM 0 15.471 1.00 34.40 0 **ATOM** 2330 CB ILE A 309 93.590 63.033 1.00 35.31 C 14.176 CG1 ILE A 309 2331 94.878 63.132 ATOM 15.004 1.00 35.26 C MOTA 2332 CG2 ILE A 309 93.709 63.974 12.980 C 1.00 34.04 MOTA 2333 CD1 ILE A 309 94.957 62.199 16.191 C 1.00 36.80 **ATOM** 2334 N CYS A 310 17.227 91.743 62.506 N 1.00 35.32 CYS A 310 **ATOM** 2335 91.380 61.483 18.209 CA 1.00 35.11 C ATOM 2336 CYS A 310 89.882 61.214 C C 18.337 1.00 34.44 CYS A 310 2337 89.072 18.449 1.00 33.70 MOTA 0 62.142 0 ATOM 2338 CYS A 310 91.922 61.873 19.587 C CB 1.00 36.64 CYS A 310 1.00 40.34 MOTA 2339 SG 91.351 60.810 20.950 S MOTA 2340 THR A 311 89.529 59.931 N 18.351 1.00 33.48 N **ATOM** THR A 311 88.143 59.500 CA 1.00 33.03 C 2341 18.487 **ATOM** 2342 C THR A 311 1.00 31.85 C 87.936 58.662 19.756 2343 86.911 19.924 1.00 32.55 ATOM THR A 311 58.016 0 MOTA 2344 58.657 THR A 311 87.720 17.287 1.00 34.00 C CB OG1 THR A 311 88.594 57.526 O MOTA 2345 17.180 1.00 36.60 ATOM C CG2 THR A 311 87.797 59.474 16.004 2346 1.00 34.76 MOTA N 2347 N THR A 312 88.912 58.667 20.650 1.00 30.55 THR A 312 88.794 C 1.00 29.18 MOTA 2348 CA 57.905 21.884 C 2349 THR A 312 87.523 22.661 MOTA C 58.238 1.00 28.00 86.831 23.144 0 THR A 312 57.344 1.00 27.47 MOTA 2350 0 58.142 C THR A 312 90.018 22.783 1.00 29.25 MOTA 2351 CB OG1 THR A 312 91.174 57.572 22.159 0 MOTA 2352 1.00 31.01 24.150 C 2353 CG2 THR A 312 89.821 57.509 1.00 30.17 MOTA 59.518 22.774 N 2354 ARG A 313 87.199 1.00 26.42 ATOM N C 86.015 23.519 2355 ARG A 313 59.899 1.00 26.82 MOTA CA C **MOTA** 1.00 27.41 C ARG A 313 84.712 59.469 22.858 2356 2357 83.699 59.237 23.531 1.00 28.22 0 MOTA ARG A 313 0 C 23.765 1.00 27.23 ARG A 313 86.025 61.407 MOTA 2358 CB

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1.00 28.90

ARG A 313

### 2360 87.183 63.318 MOTA CD ARG A 313 24.904 1.00 31.11 C MOTA 2361 NE ARG A 313 88.414 63.773 25.552 1.00 32.48 И MOTA 2362 ARG A 313 CZ25.840 88.674 65.046 1.00 32.07 C NH1 ARG A 313 87.790 ATOM 2363 1.00 30.54 65.996 25.540 N NH2 ARG A 313 ATOM 89.817 65.366 26.430 2364 1.00 31.87 N VAL A 314 84.746 59.339 1.00 26.45 MOTA 2365 N 21.540 N VAL A 314 83.576 20.774 MOTA 2366 58.952 C CA 1.00 25.00 2367 **VAL A 314** 57.446 20.698 C 83.364 $\mathbf{C}$ MOTA 1.00 24.59 **ATOM** 2368 VAL A 314 82.253 56.963 20.890 0 1.00 24.29 0 59.496 ATOM 2369 VAL A 314 83.672 19.338 CB C 1.00 26.31 CG1 VAL A 314 MOTA 2370 82.429 59.127 18.558 1.00 26.03 C 2371 CG2 VAL A 314 60.998 C 19.371 MOTA 83.866 1.00 25.74 MOTA 2372 VAL A 315 84.428 56.704 20.416 N 1.00 24.12 N 2373 VAL A 315 **ATOM** 84.322 55.255 20.286 C CA 1.00 24.14 VAL A 315 MOTA 2374 C 54.460 21.595 1.00 23.17 C 84.419 2375 **VAL A 315** 83.837 53.387 21.713 0 1.00 22.34 MOTA 2376 VAL A 315 85.351 54.737 19.266 1.00 23.99 CB C MOTA CG1 VAL A 315 55.352 C MOTA 85.058 17.897 1.00 24.71 2377 **ATOM** 2378 CG2 VAL A 315 86.754 55.099 19.705 C 1.00 25.41 ALA A 316 85.137 54.992 22.577 N MOTA 2379 N 1.00 22.80 2380 ALA A 316 85.273 54.322 23.864 1.00 22.89 C CA MOTA ALA A 316 84.453 55.075 C C MOTA 2381 24.907 1.00 23.76 2382 ALA A 316 84.096 54.525 25.955 1.00 23.93 0 0 MOTA 86.739 54.277 24.278 **ATOM** 2383 ALA A 316 C CB 1.00 21.84 **GLY A 317** 56.337 24.607 N MOTA 2384 N 84.154 1.00 24.54 2385 83.378 1.00 24.66 C MOTA CA **GLY A 317** 57.167 25.511 26.744 **GLY A 317** 84.154 57.582 1.00 25.96 C **ATOM** 2386 C **GLY A 317** 83.558 57.954 1.00 26.42 0 MOTA 2387 27.759 VAL A 318 85.482 57.545 26.646 1.00 25.51 MOTA 2388 N N 27.760 C 2389 86.369 57.879 MOTA CA VAL A 318 1.00 24.79 87.219 2390 C VAL A 318 27.554 1.00 25.38 MOTA 59.133 87.633 59.437 26.438 2391 VAL A 318 1.00 25.50 MOTA 0 87.317 CB VAL A 318 56.693 28.049 1.00 24.92 MOTA 2392 C CG1 VAL A 318 29.090 MOTA 2393 88.363 57.081 1.00 24.67 C 86.507 55.500 28.535 1.00 23.44 2394 CG2 VAL A 318 MOTA 87.477 59.851 N 2395 N GLY A 319 28.645 1.00 24.73 MOTA CA GLY A 319 28.578 C 2396 88.304 61.043 1.00 25.39 ATOM 87.980 62.097 2397 GLY A 319 29.620 1.00 25.66 MOTA C 87.076 61.926 1.00 25.40 30.437 2398 GLY A 319 MOTA 29.609 N **ATOM** 2399 VAL A 320 88.737 63.188 1.00 24.72 N C 88.484 64.281 30.535 1.00 25.55 VAL A 320 **ATOM** 2400 CA $\mathbf{C}$ 65.581 29.903 88.944 1.00 23.92 VAL A 320 ATOM 2401 C 0 65.653 29.331 1.00 22.93 2402 90.033 MOTA 0 VAL A 320 C 89.203 64.087 31.899 1.00 27.29 ATOM CB VAL A 320 2403 1.00 32.20 31.693 CG1 VAL A 320 90.667 64.017 ATOM 2404 C 65.241 32.838 1.00 26.56 CG2 VAL A 320 88.884 ATOM 2405 88.100 66.625 29.980 PRO A 321 1.00 23.64 2406 N ATOM

### ATOM 2407 CA PRO A 321 29.408 88.438 67.928 1.00 22.67 C **ATOM** 2408 PRO A 321 89.872 C 68.279 29.791 1.00 22.52 C 2409 PRO A 321 90.232 **ATOM** 68.263 0 30.961 1.00 23.18 0 87.381 68.823 ATOM 2410 CB PRO A 321 30.028 C 1.00 23.67 ATOM 2411 CG PRO A 321 86.164 67.896 29.986 C 1.00 20.93 2412 PRO A 321 **ATOM** 30.640 CD 86.780 66.686 1.00 22.53 C **ATOM GLN A 322** 90.679 2413 68.588 1.00 22.16 N 28.786 N ATOM 2414 **GLN A 322** 92.099 CA 28.948 1.00 21.62 C 68.880 MOTA 2415 C **GLN A 322** 92.573 69.884 30.002 1.00 20.84 C 69.609 ATOM 2416 **GLN A 322** 93.551 1.00 21.63 0 30.679 0 **GLN A 322** MOTA 92.684 69.245 2417 CB C 27.578 1.00 22.97 2418 **GLN A 322** 94.217 69.292 **ATOM** CG 27.491 1.00 24.38 C **ATOM** 2419 CD **GLN A 322** 1.00 23.75 94.896 67.951 27.725 C OE1 GLN A 322 **ATOM** 96.067 67.776 27.390 2420 1.00 26.82 0 2421 NE2 GLN A 322 ATOM 94.176 67.011 28.314 1.00 23.57 N 2422 **VAL A 323** 91.932 71.037 30.151 ATOM N 1.00 19.60 N VAL A 323 92.401 71.970 **ATOM** 2423 CA 31.175 1.00 19.52 C **ATOM** 2424 **VAL A 323** C 32.572 92.280 71.357 C 1.00 20.46 MOTA 2425 **VAL A 323** 1.00 22.03 0 93.144 71.556 33.420 0 **VAL A 323** 2426 CB MOTA 91.645 73.322 C 31.143 1.00 17.80 2427 CG1 VAL A 323 MOTA 92.026 74.177 32.359 C 1.00 17.57 2428 CG2 VAL A 323 92.009 74.070 C MOTA 29.888 1.00 18.00 THR A 324 91.210 70.616 2429 1.00 21.51 MOTA Ν 32.815 N **ATOM** 2430 CA THR A 324 91.029 69.961 34.106 1.00 21.74 C THR A 324 MOTA C 1.00 22.09 2431 92.056 68.845 C 34.307 THR A 324 35.393 1.00 22.04 MOTA 2432 0 92.628 68.704 0 MOTA 69.383 34.221 2433 CB THR A 324 89.616 1.00 21.42 C OG1 THR A 324 34.469 MOTA 88.696 2434 70.448 1.00 23.37 0 THR A 324 **ATOM** 2435 89.532 68.370 35.335 C CG2 1.00 24.31 33.256 **ATOM** 92.296 68.062 2436 ALA A 325 N 1.00 21.63 N 93.253 ATOM 2437 CA ALA A 325 66.958 33.324 1.00 23.09 C 2438 ALA A 325 94.669 33.668 C ATOM C 67.437 1.00 23.65 2439 95.325 66.865 ATOM О ALA A 325 34.540 1.00 23.99 0 C MOTA 2440 66.192 ALA A 325 93.262 32.006 1.00 22.44 CBN ILE A 326 MOTA 2441 $\mathbf{N}$ 95.134 68.472 32.967 1.00 23.50 ILE A 326 96.458 69.067 C MOTA 2442 CA 33.182 1.00 23.40 C 2443 ILE A 326 MOTA C 96.576 69.483 34.641 1.00 24.71 MOTA 2444 97.520 69.124 0 ILE A 326 35.337 1.00 24.66 0 C ILE A 326 1.00 22.70 96.644 70.329 32.295 MOTA 2445 СВ C CG1 ILE A 326 96.680 69.913 30.821 1.00 22.18 MOTA 2446 32.699 C CG2 ILE A 326 **ATOM** 2447 97.900 71.096 1.00 20.32 1.00 22.52 C 71.078 ATOM 2448 CD1 ILE A 326 96.689 29.861 95.589 N 2449 70.244 35.082 1.00 25.51 ATOM TYR A 327 N 1.00 27.53 C TYR A 327 70.739 MOTA 2450 CA 95.512 36.439 C 95.493 TYR A 327 69.644 37.506 1.00 26.67 ATOM 2451 C 69.706 38.470 2452 TYR A 327 0 96.234 1.00 26.34 ATOM 0 C 94.278 71.615 36.527 1.00 31.94 ATOM 2453 CBTYR A 327

### ATOM 2454 CG TYR A 327 93.867 72.043 37.900 1.00 36.64 C CD1 TYR A 327 MOTA 71.127 38.812 2455 93.354 1.00 40.23 C MOTA CD2 TYR A 327 2456 93.868 73.375 38.243 1.00 38.78 CE1 TYR A 327 92.839 71.536 ATOM 2457 40.027 1.00 42.47 C TYR A 327 ATOM 2458 CE2 93.358 73.790 C 39.431 1.00 42.01 CZ92.838 72.873 MOTA **TYR A 327** 1.00 43.22 2459 40.329 C MOTA TYR A 327 92.305 73.316 41.524 2460 0 OH 1.00 47.71 **ATOM** 2461 **ASP A 328** 94.635 68.650 1.00 27.41 И 37.347 N ATOM 2462 **ASP A 328** 94.573 67.559 38.312 C CA 1.00 28.23 66.825 38.383 **ATOM** 2463 C **ASP A 328** 95.910 1.00 28.97 **ASP A 328 ATOM** 2464 96.342 66.414 39.455 1.00 30.49 0 **ASP A 328** MOTA 2465 CB 93.469 66.570 37.925 1.00 28.63 C **ASP A 328** 38.042 92.074 1.00 29.55 C MOTA 2466 CG 67.171 OD1 ASP A 328 37.643 91.096 66.496 0 ATOM 2467 1.00 28.91 OD2 ASP A 328 **MOTA** 2468 91.954 68.312 38.541 1.00 30.66 0 **ALA A 329** 66.666 **ATOM** 2469 N 96.568 37.237 1.00 28.70 N ALA A 329 2470 97.848 65.969 37.185 C MOTA CA 1.00 27.96 2471 **ALA A 329** 98.993 66.816 37.713 1.00 27.90 C MOTA C 2472 ALA A 329 99.898 0 ATOM 66.303 38.368 1.00 27.58 ALA A 329 C MOTA 2473 CB 98.150 65.534 35.755 1.00 28.88 **ALA A 330** 68.112 37.417 98.950 MOTA 2474 N 1.00 27.54 Ν MOTA ALA A 330 69.041 37.839 C 2475 CA 99.993 1.00 27.36 2476 **ALA A 330** 69.195 39.352 1.00 27.62 C C 100.035 MOTA 2477 ALA A 330 101.039 69.631 39.909 0 0 1.00 27.21 MOTA 99.784 70.397 1.00 26.61 C MOTA 2478 CB ALA A 330 37.175 ALA A 331 98.941 68.841 40.016 N 1.00 28.38 MOTA 2479 Ν ALA A 331 41.467 C ATOM 2480 CA98.882 68.923 1.00 29.60 ALA A 331 99.870 67.902 42.005 C 2481 MOTA C 1.00 30.94 68.144 42.995 1.00 31.96 100.556 MOTA 2482 О ALA A 331 68.601 41.955 1.00 29.46 CB ALA A 331 97.489 MOTA 2483 2484 99.941 MOTA 66.756 41.334 VAL A 332 1.00 31.86 VAL A 332 100.851 41.723 1.00 31.71 65.683 ATOM 2485 CAC MOTA 2486 VAL A 332 102.276 65.984 41.267 1.00 32.27 VAL A 332 103.231 41.981 1.00 32.86 O MOTA 65.700 2487 C CB VAL A 332 MOTA 2488 100.408 64.328 41.122 1.00 30.47 41.512 1.00 29.17 C CG1 VAL A 332 101.394 63.229 MOTA 2489 C 99.009 41.602 ATOM CG2 VAL A 332 63.989 1.00 29.70 2490 N 66.561 40.078 ALA A 333 1.00 32.53 MOTA 102.412 2491 N C 103.723 66.889 39.543 1.00 33.43 CA ALA A 333 MOTA 2492 C 104.469 1.00 35.24 ALA A 333 67.763 40.528 MOTA 2493 C 0 ALA A 333 105.677 67.623 40.701 1.00 35.30 **ATOM** 2494 0 C 38.218 ALA A 333 103.587 1.00 32.68 67.607 ATOM 2495 CBN 68.675 41.178 1.00 36.59 ARG A 334 103.757 2496 MOTA N 42.136 C 1.00 38.67 104.422 69.544 MOTA 2497 CA ARG A 334 C 104.682 68.857 ARG A 334 43.458 1.00 38.64 ATOM 2498 0 1.00 39.25 **ATOM** 2499 105.646 69.181 44.140 0 ARG A 334 C 103.650 70.853 42.343 1.00 39.27 ARG A 334 ATOM 2500 CB

MOTA	2501	CG	ARG	A	334	102.152	70.742	42.460	1.00	41.57	C
MOTA	2502	CD	ARG	A	334	101.611	72.112	42.821	1.00	42.79	C
MOTA	2503	NE	ARG	Α	334	102.174	73.143	41.953	1.00	41.68	N
MOTA	2504	CZ	ARG	Α	334	102.177	74.441	42.245	1.00	41.25	C
MOTA	2505	NH1	ARG	A	334	101.648	74.876	43.384	1.00	39.55	N
MOTA	2506	NH2	ARG	A	334	102.728	75.304	41.405	1.00	40.34	N
MOTA	2507	Ŋ	GLU	A	335	103.833	67.902	43.814	1.00	38.96	N
MOTA	2508	CA	GLU	Α	335	104.029	67.158	45.046	1.00	39.04	C
ATOM	2509	C	GLU	A	335	105.352	66.414	44.911		37.56	C
ATOM	2510	0	GLU	A	335	106.158	66.383	45.836			0
ATOM	2511	CB	GLU			102.895	66.157	45.258	1.00		C
MOTA	2512	CG	GLU			103.154	65.182	46.393	1.00		C
MOTA	2513	CD	GLU			102.064	64.136	46.538	1.00		C
ATOM	2514	OE1	GLU			102.219	63.227	47.383	1.00		0
MOTA	2515	OE2	GLU	A	335	101.048	64.224	45.812	1.00		0
MOTA	2516	N	TYR	A	336	105.571	65.828	43.741	1.00	35.56	N
MOTA	2517	CA	TYR	A	336	106.786	65.075	43.458	1.00	33.65	C
MOTA	2518	C	TYR			107.915	65.956	42.950	1.00	32.06	C
MOTA	2519	0	TYR	A	336	109.037	65.493	42.753	1.00	31.27	0
MOTA	2520	CB	TYR			106.508	63.979	42.418	1.00	34.56	C
MOTA	2521	CG	TYR	A	336	105.700	62.806	42.928	1.00		C
MOTA	2522	CD1	TYR	A	336	106.183	61.504	42.802	1.00	35.01	C
ATOM	2523	CD2	TYR	A	336	104.465	62.992	43.544	1.00	35.65	C
MOTA	2524	CE1	TYR	A	336	105.460	60.418	43.278	1.00	35.67	C
MOTA	2525	CE2	TYR			103.730	61.910	44.025	1.00	36.28	C
MOTA	2526	CZ	TYR	A	336	104.234	60.624	43.890	1.00		C
MOTA	2527	OH	TYR	Α	336	103.516	59.548	44.372		36.21	0
ATOM	2528	N	GLY			107.619	67.230	42.738		31.22	N
MOTA	2529	CA	GLY		-	108.635	68.122	42.229	1.00	29.58	C
MOTA	2530	C	GLY			109.068	67.679	40.847	1.00		C
MOTA	2531	0	GLY			110.232	67.802	40.488	1.00		0
MOTA	2532	N	LYS	A	338	108.128	67.142	40.073	1.00		N
ATOM	2533	CA	LYS	A	338	108.413	66.698	38.713	1.00		C
ATOM	2534	C	LYS			107.596	67.477	37.683	1.00		C
MOTA	2535	0	LYS	Α	338	106.830	68.377	38.040		27.37	0
MOTA	2536	CB			338	108.180	65.191	38.585	1.00		C
ATOM	2537	CG	LYS	Α	338	109.235	64.382	39.330	1.00		C
MOTA	2538	CD	LYS	A	338	109.034	62.886	39.194	1.00		C
MOTA	2539	CE	LYS	A	338	110.170	62.116	39.867	1.00	38.61	C
ATOM	2540	NZ	LYS	Α	338	110.001	60.628	39.761	1.00		N
MOTA	2541	N	THR	Α	339	107.749	67.131	36.407	1.00		N
MOTA	2542	CA	THR	A	339	107.056	67.863	35.356	1.00		C
MOTA	2543	C	THR	A	339	105.980	67.165	34.517		26.06	C
MOTA	2544	0	THR	A	339	105.806	65.943	34.542		25.01	0
ATOM	2545	CB	THR	A	339	108.072	68.484	34.409		26.34	C
MOTA	2546	OG1	THR	A	339	108.901	67.454	33.864		25.90	0
MOTA	2547	CG2	THR	A	339	108.934	69.484	35.155	1.00	26.42	C

ATOM	2548	N	ILE	A	340	105.267	67.988	33.757	1.00 25.60	N
ATOM	2549	CA	ILE	A	340	104.174	67.543	32.911	1.00 24.59	C
MOTA	2550	С	ILE	A	340	104.152	68.189	31.525	1.00 23.54	С
MOTA	2551	0	ILE	A	340	104.555	69.337	31.342	1.00 21.85	0
MOTA	2552	CB	ILE	Α	340	102.842	67.794	33.644	1.00 25.00	
ATOM	2553	CG1			340	102.598	66.631	34.599	1.00 26.20	
ATOM	2554	CG2				101.707	68.034	32.668	1.00 25.49	
ATOM	2555	CD1		A		101.382	66.774	35.436	1.00 29.56	
ATOM	2556	N	ILE	A	341	103.695	67.414	30.548	1.00 22.21	
ATOM	2557	CA				103.573	67.873	29.176	1.00 20.48	
ATOM	2558	C				102.080	67.824	28.863	1.00 19.91	
ATOM	2559	0		A		101.439	66.793	29.068	1.00 18.38	
ATOM	2560	CB		A	341	104.346	66.941	28.210	1.00 21.69	
ATOM	2561	CG1		A	341	105.849	67.038	28.496	1.00 23.08	
ATOM	2562	CG2		A		104.043	67.290	26.755	1.00 19.22	
ATOM	2563	CD1		A	341	106.695	66.098	27.657	1.00 23.45	
ATOM	2564	N	ALA ALA			101.530 100.121	68.958 69.060	28.427 28.046	1.00 19.15 1.00 19.91	
ATOM	2565	CA C	ALA			100.121	68.553	26.601	1.00 19.91	
ATOM ATOM	2566 2567	0	ALA			100.556	69.205	25.680	1.00 20.83	
ATOM	2568	CB	ALA			99.663	70.504	28.125	1.00 20.43	
ATOM	2569	N	ASP			99.451	67.388	26.418	1.00 21.90	
ATOM	2570	CA	ASP			99.390	66.729	25.121	1.00 22.21	
ATOM	2571	C	ASP			98.038	66.728	24.406	1.00 23.22	
ATOM	2572	0	ASP			97.128	66.005	24.796	1.00 23.89	
ATOM	2573	СВ	ASP			99.863	65.283	25.318	1.00 23.79	
ATOM	2574	CG	ASP			99.944	64.499	24.028	1.00 26.58	
ATOM	2575	OD1	ASP			100.061	63.251	24.106	1.00 25.81	
ATOM	2576	OD2	ASP			99.911	65.122	22.941	1.00 27.87	
ATOM	2577	N	GLY	A	344	97.917	67.541	23.359	1.00 24.09	N
ATOM	2578	CA	GLY	Α	344	96.696	67.569	22.565	1.00 24.48	C
ATOM	2579	С	GLY	Α	344	95.592	68.587	22.818	1.00 24.98	C
MOTA	2580	0	GLY	Α	344	95.510	69.210	23.888	1.00 25.91	. 0
ATOM	2581	N	GLY	A	345	94.737	68.753	21.809	1.00 23.62	N
MOTA	2582	CA	GLY	Α	345	93.614	69.664	21.922	1.00 22.65	C
MOTA	2583	С	GLY	A	345	93.944	71.128	21.758	1.00 22.90	C
ATOM	2584	0	GLY	A	345	93.060	71.969	21.869	1.00 23.02	0
MOTA	2585	N	ILE	Α	346	95.207	71.443	21.498	1.00 22.89	
ATOM	2586	CA	ILE	A	346	95.637	72.827	21.322	1.00 23.44	
MOTA	2587	C	ILE	Α	346	95.528	73.267	19.869	1.00 25.23	
ATOM	2588	0	ILE	A	346	96.157	72.681	18.992	1.00 26.62	
MOTA	2589	CB			346	97.090	73.005	21.774	1.00 22.40	
ATOM	2590	CG1			346	97.160	72.887	23.297	1.00 23.15	
ATOM	2591		ILE			97.631	74.327	21.268	1.00 22.29	
ATOM	2592	CD1			346	98.558	72.897	23.857	1.00 25.18	
ATOM	2593	N			347	94.744	74.305	19.603	1.00 25.74	
ATOM	2594	CA	LYS	A	347	94.605	74.756	18.228	1.00 26.93	C C

ATOM	2595	C	LYS	A	347	95.037	76.200	18.004	1.00	26.46	C
ATOM	2596	0	LYS	A	347	95.310	76.602	16.874	1.00	27.31	0
MOTA	2597	CB	LYS	A	347	93.169	74.501	17.744	1.00	29.47	C
MOTA	2598	CG	LYS	Α	347	92.093	75.047	18.644	1.00	32.60	C
ATOM	2599	CD			347	90.778	74.293	18.454	1.00	36.26	C
ATOM	2600	CE			347	89.680	74.859	19.374	1.00	39.81	C
ATOM	2601	NZ	LYS			90.034	74.868	20.845	1.00		N
ATOM	2602	N	TYR			95.136	76.968	19.081	1.00	24.97	N
ATOM	2603	CA	TYR			95.552	78.357	18.982	1.00		C
ATOM	2604	C	TYR			96.692	78.637	19.951	1.00		С
ATOM	2605	0	TYR			96.810	77.976	20.987	1.00		0
ATOM	2606	CB	TYR			94.384	79.280	19.313	1.00	25.92	C
ATOM	2607	CG	TYR			93.224	79.180	18.362	1.00		C
ATOM	2608	CD1	TYR			93.363	79.544	17.014	1.00	31.01	C
ATOM	2609	CD2	TYR			91.975	78.759	18.806	1.00	30.18	C
ATOM	2610	CE1	TYR			92.280	79.496	16.139	1.00	31.84	C
ATOM	2611	CE2			348	90.886	78.706	17.940	1.00	32.70	C
ATOM	2612	CZ	TYR			91.046	79.078	16.614	1.00	33.71	C
ATOM	2613	OH	TYR		348	89.960	79.054	15.775	1.00	37.15	O N
ATOM	2614	N	SER			97.523	79.626 79.970	19.629	1.00	23.37	N
ATOM	2615 2616	CA C	SER SER			98.632 98.093	80.308	20.503 21.894	1.00	23.44	C
ATOM		0	SER			98.747	80.049	22.900	1.00		0
ATOM	2617 2618	СВ	SER			99.432	81.151	19.936		23.65	C
ATOM ATOM	2619	OG	SER			98.657	82.331	19.874	1.00	24.07	0
ATOM	2620	N	GLY			96.896	80.880	21.950	1.00		N
ATOM	2621	CA	GLY			96.310	81.212	23.234	1.00		C
ATOM	2622	C	GLY			96.093	79.964	24.071	1.00		Ċ
ATOM	2623	0	GLY			96.186	80.009	25.291	1.00	23.00	0
ATOM	2624	N	ASP			95.798	78.847	23.407	1.00		N
ATOM	2625	CA	ASP			95.573	77.573	24.085	1.00		C
ATOM	2626	C	ASP			96.850	77.093	24.770	1.00		C
ATOM	2627	0				96.798	76.434	25.811	1.00		0
ATOM	2628	СВ	ASP			95.097	76.509	23.088		25.03	C
ATOM	2629	CG	ASP			93.660	76.720	22.638		27.18	C
ATOM	2630	OD1	ASP		351	93.215	76.016	21.707	1.00		0
ATOM	2631	OD2			351	92.966	77.576	23.226	1.00		0
ATOM	2632	N			352	97.994	77.421	24.178	1.00		N
ATOM	2633	CA			352	99.274	77.032	24.749	1.00	20.34	С
ATOM	2634	С	ILE	Α	352	99.452	77.716	26.102	1.00	21.36	C
ATOM	2635	0	ILE	Α	352	99.879	77.084	27.069	1.00	22.39	0
ATOM	2636	CB	ILE	Α	352	100.445	77.431	23.830	1.00	18.46	C
ATOM	2637	CG1	ILE	A	352	100.296	76.753	22.474	1.00	17.19	С
MOTA	2638	CG2	ILE	A	352	101.756	77.042	24.462	1.00	17.65	C
ATOM	2639	CD1	ILE	A	352	101.395	77.086	21.512	1.00	17.04	C
MOTA	2640	N	VAL	A	353	99.122	79.005	26.172	1.00	20.35	N
MOTA	2641	CA	VAL	A	353	99.269	79.753	27.413	1.00	20.32	C

ATO ATO ATO ATO ATO ATO ATO ATO ATO ATO	OM 26 OM 26 OM 26 OM 26 OM 26 OM 26	542 C 543 O 544 CE 545 CG 546 CG	VAL VAL VAL	A A	353 353 353	98.328 98.671 98.997	79.228 79.204 81.250	28.496 29.671 27.189	1.00 19.94 1.00 20.43 1.00 21.35	C 0 C
ATO	OM 26 OM 26 OM 26 OM 26 OM 26	544 CE 545 CG 546 CG	VAL	A	353					
ATO	OM 26 OM 26 OM 26	545 CG 546 CG	1 VAL			98.997	81.250	27.189	1.00 21.35	C
ATO	OM 26 OM 26 OM 26	646 CG		Α	252					
ATO	OM 26		:2 WAT.		333	99.315	82.026	28.455	1.00 21.91	С
ATO ATO ATO ATO ATO ATO ATO ATO ATO	OM 26	547 N	, v F111	Α	353	99.826	81.765	26.024	1.00 22.51	C
ATO		, _ ,	LYS	Α	354	97.131	78.809	28.098	1.00 21.28	N
ATO ATO ATO ATO ATO ATO ATO ATO	OM 24	48 CA	LYS	A	354	96.168	78.266	29.049	1.00 20.46	C
ATO ATO ATO ATO ATO ATO ATO ATO	Or1 20	649 C	LYS	Α	354	96.674	76.925	29.563	1.00 20.59	C
ATO ATO ATO ATO ATO ATO ATO	OM 26	50 O	LYS	Α	354	96.587	76.640	30.758	1.00 21.00	0
ATO ATO ATO ATO ATO ATO ATO	OM 26	551 CE	LYS	A	354	94.797	78.067	28.388	1.00 21.23	C
ATO ATO ATO ATO ATO ATO ATO	OM 26	552 CG	LYS	A	354	94.090	79.344	27.951	1.00 20.48	С
ATO ATO ATO ATO ATO ATO	OM 26	553 CE	LYS	A	354	92.766	79.032	27.264	1.00 19.71	C
ATO ATO ATO ATO ATO	OM 26	554 CE	LYS	Α	354	92.118	80.297	26.720	1.00 22.52	С
ATO ATO ATO ATO ATO	OM 26	555 NZ	LYS	Α	354	90.917	80.013	25.879	1.00 21.12	N
ATO ATO ATO ATO	OM 26	556 N	ALA	Α	355	97.191	76.099	28.656	1.00 19.85	N
ATO ATO ATO	OM 26	557 CA	ALA	A	355	97.716	74.782	29.017	1.00 20.80	C
ATO ATO	OM 26	558 C	ALA	A	355	98.886	74.933	29.978	1.00 21.77	C
OTA OTA	OM 26	559 0	ALA	Α	355	99.037	74.146	30.908	1.00 22.61	0
ATO	OM 26	60 CE			355	98.163	74.022	27.773	1.00 19.41	C
	OM 26	61 N			356	99.722	75.940	29.753	1.00 21.62	Ŋ
		62 CA			356	100.858	76.168	30.630	1.00 22.27	C
ATO		63 C			356	100.367	76.727	31.957	1.00 23.40	C
ATO		664 0			356	100.846	76.342	33.015	1.00 24.73	0
ATO		65 CE			356	101.843	77.145	29.986	1.00 21.42	C
AT(		66 CG			356	102.518	76.717	28.684	1.00 20.49	C
AT(		67 CD			356 356	103.328	77.875	28.141	1.00 21.00	C
AT(		68 CD			356	103.386 99.396	75.488 77.629	28.925 31.900	1.00 21.89 1.00 23.83	N
AT(		569 N 570 CA			357 357	98.858	78.243	33.109	1.00 24.49	C
ATO ATO		570 CA			357	98.115	77.252	34.002	1.00 24.43	C
AT(		572 O			357	98.034	77.445	35.210	1.00 24.39	ō
ATO		773 CE			357	97.943	79.396	32.739	1.00 25.49	C
AT		574 N			358	97.577	76.191	33.412	1.00 23.94	N
ATO		75 CA			358	96.856	75.192	34.189	1.00 23.85	C
ATO		576 C			358	97.818	74.209	34.853	1.00 24.49	C
ATO		577 0	ALA	Α	358	97.391	73.320	35.595	1.00 24.74	0
ATO	OM 26	78 CE	ALA	Α	358	95.865	74.440	33.300	1.00 24.65	C
ATO	OM 26	579 N	GLY	A	359	99.113	74.349	34.579	1.00 24.04	N
ATO	OM 26	80 CA	GLY	Α	359	100.074	73.460	35.206	1.00 23.58	C
AT(	OM 26	581 C	GLY	A	359	101.065	72.763	34.300	1.00 23.49	C
ATO	OM 26	82 0	GLY	A	359	101.964	72.073	34.786	1.00 24.97	0
ATO	OM 26	83 N	GLY	Α	360	100.924	72.929	32.990	1.00 22.19	N
ATO		584 CP			360	101.847	72.279	32.083	1.00 19.65	C
AT(		85 C			360	103.209	72.936	32.105	1.00 20.75	C
ATO	OM 26	586 O			360	103.312	74.134	32.324	1.00 20.45	0
AT(			አ ሮእ፣	7	361	104.258	72.148	31.887	1.00 20.92	N
ATO		587 N 588 CA			361	105.630	72.655	31.855	1.00 22.23	C

# TABLE 7 ATOM 26

ATOM	2689	C	ASN	A	361	106.051	72.782	30.402	1.00	22.46	C
ATOM	2690	0	ASN	A	361	107.060	73.407	30.079	1.00	22.58	0
ATOM	2691	CB	ASN	A	361	106.568	71.697	32.589	1.00	22.17	C
ATOM	2692	CG	ASN	A	361	106.338	71.703	34.085	1.00	25.53	C
MOTA	2693	OD1	ASN	A	361	106.646	72.688	34.763	1.00	27.16	0
MOTA	2694	ND2	ASN	A	361	105.766	70.624	34.607	1.00	25.19	N
MOTA	2695	N	ALA	A	362	105.251	72.178	29.530	1.00	22.54	N
MOTA	2696	CA	ALA	A	362	105.481	72.193	28.095	1.00	21.37	C
MOTA	2697	C	ALA	A	362	104.229	71.625	27.432	1.00	21.05	C
MOTA	2698	0	ALA	A	362	103.423	70.969	28.084	1.00	18.71	0
MOTA	2699	CB	ALA			106.704	71.341	27.753		20.68	С
MOTA	2700	N	VAL			104.058	71.895	26.142		22.66	N
MOTA	2701	CA	VAL			102.903	71.377	25.410		22.68	С
MOTA	2702	C	VAL			103.389	70.579	24.210		22.35	C
MOTA	2703	0	VAL			104.423	70.893	23.638		23.37	0
ATOM	2704	CB	VAL			101.964	72.521	24.910		22.34	C
ATOM	2705	CG1	VAL			101.440	73.333	26.088		21.37	C
ATOM	2706	CG2	VAL		363	102.702	73.418	23.929		22.84	C
HETATM	2707	N	MSE			102.662	69.530	23.851		21.88	N
	2708	CA	MSE			103.027	68.726	22.692		22.51	C
	2709	C			364	102.014	69.063	21.608		23.49	C
HETATM		0	MSE		364	100.794	69.022	21.832		23.44	0
HETATM		CB	MSE			103.011	67.231	23.028		24.05	C
HETATM		CG	MSE			103.223	66.330	21.821 22.241		27.31 35.51	SE
HETATM HETATM	2713	SE CE	MSE MSE			103.374 105.096	64.556 64.508	22.891		32.69	C
ATOM	2715	N			365	102.531	69.420	20.436		23.87	N
ATOM	2716	CA	LEU			101.704	69.831	19.315		24.27	C
ATOM	2717	C			365	101.822	68.920	18.108		25.78	Č
ATOM	2718	0	LEU			102.926	68.614	17.666		26.09	0
MOTA	2719	CB			365	102.104	71.238	18.905		22.73	C
ATOM	2720	CG			365	102.093	72.219	20.060		23.17	C
ATOM	2721	CD1	LEU			102.705	73.527	19.616		24.46	C
ATOM	2722	CD2	LEU	Α	365	100.670	72.378	20.574	1.00	22.59	C
ATOM	2723	N	GLY	Α	366	100.678	68.508	17.569	1.00	26.69	И
ATOM	2724	CA	GLY	Α	366	100.667	67.651	16.398	1.00	28.30	C
ATOM	2725	C	GLY	Α	366	100.045	68.365	15.213	1.00	29.57	С
ATOM	2726	0	GLY	A	366	100.717	68.663	14.226	1.00	29.34	0
MOTA	2727	N	SER	Α	367	98.753	68.644	15.327	1.00	30.92	N
MOTA	2728	CA	SER	Α	367	97.993	69.341	14.298	1.00	33.00	C
ATOM	2729	C	SER	A	367	98.648	70.637	13.803	1.00	34.00	С
ATOM	2730	0	SER	A	367	98.750	70.848	12.598		33.51	0
MOTA	2731	CB	SER	A	367	96.590	69.646	14.832	1.00	34.28	C
MOTA	2732	OG	SER	A	367	95.800	70.345	13.885		34.75	0
HETATM	2733	N	MSE	A	368	99.097	71.501	14.713		34.95	N
HETATM	2734	CA	MSE	A	368	99.712	72.765	14.296		37.45	C
HETATM	2735	С	MSE	A	368	100.963	72.668	13.424	1.00	36.66	С

HETATM	2736	0	MSE	A	368	101.316	73.635	12.748	1.00	36.99	0
HETATM	2737	CB	MSE	A	368	100.000	73.664	15.506		40.56	С
HETATM	2738	CG	MSE	A	368	98.719	74.111	16.191		48.02	С
HETATM	2739	SE	MSE	A	368	98.916	75.343	17.492		56.80	SE
HETATM	2740	CE	MSE	A	368	99.964	74.527	18.572		55.31	С
MOTA	2741	N	PHE	A	369	101.626	71.513	13.417	1.00	34.82	N
MOTA	2742	CA	PHE	A	369	102.830	71.339	12.603		31.76	С
MOTA	2743	C	PHE	A	369	102.600	70.421	11.416	1.00	31.43	С
ATOM	2744	0	PHE	A	369	103.378	70.425	10.467	1.00	30.91	0
MOTA	2745	CB	PHE	A	369	103.977	70.786	13.460		29.79	С
MOTA	2746	CG	PHE	Α	369	104.507	71.765	14.473		28.59	С
ATOM	2747	CD1	PHE	A	369	104.387	71.518	15.829		27.53	C
MOTA	2748	CD2	PHE	Α	369	105.121	72.947	14.062		27.73	С
MOTA	2749	CE1	PHE	A	369	104.872	72.444	16.771		27.77	С
MOTA	2750	CE2	PHE	A	369	105.605	73.874	14.990		27.15	C
MOTA	2751	CZ	PHE	A	369	105.481	73.622	16.345	1.00	26.83	С
MOTA	2752	N	ALA	A	370	101.510	69.658	11.479	1.00	32.61	N
ATOM	2753	CA	ALA	A	370	101.108	68.668	10.465	1.00	33.73	C
MOTA	2754	C	ALA	Α	370	101.243	68.936	8.953		34.56	C
MOTA	2755	0	ALA	A	370	101.405	67.982	8.173	1.00	35.53	0
MOTA	2756	CB	ALA	A	370	99.682	68.213	10.761	1.00	33.78	С
MOTA	275 <b>7</b>	N	GLY	A	371	101.165	70.191	8.518		33.99	N
MOTA	2758	CA	GLY	A	371	101.282	70.453	7.090		33.01	С
MOTA	2759	C	GLY	A	371	102.500	71.252	6.666		33.01	С
ATOM	2760	0	GLY	A	371	102.537	71.811	5.564		32.73	0
MOTA	2761	N	THR	A	372	103.508	71.299	7.530		32.22	N
MOTA	2762	CA	THR	A	372	104.724	72.055	7.244		32.27	C
MOTA	2763	C			372	105.688	71.363	6.276		33.94	C
MOTA	2764	0			372	105.448	70.244	5.828		35.02	0
MOTA	2765	CB			372	105.479	72.426	8.567		30.26	C
MOTA	2766	OG1			372	105.828	71.242	9.298		27.73	0
ATOM	2767	CG2			372	104.601	73.299	9.440		28.61	C
MOTA	2768	N			373	106.771	72.052	5.937		35.72	N
MOTA	2769	CA			373	107.775	71.506	5.040		37.55	C
ATOM	2770	C			373	108.346	70.234	5.665		39.10	C
ATOM	2771	0			373	108.423	69.187	5.023		39.87	0
ATOM	2772	CB			373	108.893	72.531	4.844		39.43	C
ATOM	2773	ÇG			373	108.394	73.826	4.223		42.12	C
ATOM	2774	OD1			373	109.121	74.847	4.253		42.31	0
ATOM	2775	OD2			373	107.267	73.815	3.687		45.19	0
MOTA	2776	N			374	108.720	70.334	6.937		39.77	N
MOTA	2777	CA			374	109.312	69.224	7.676		40.52	C
MOTA	2778	C			374	108.430	67.991	7.949		40.61	C
MOTA	2779	0			374	108.938	66.870	7.961		41.45	0
MOTA	2780	CB			374	109.859	69.740	9.003		40.97	C
MOTA	2781	CG			374	110.940	70.800	8.893		43.47	C
MOTA	2782	CD	GLU	A	374	110.498	72.042	8.138	1.00	46.43	С

MOTA	2783	OE1	GLU	A	374	109.291	72.384	8.186	1.00	46.56	0
MOTA	2784	OE2	GLU	Α	374	111.365	72.704	7.520	1.00	48.05	0
MOTA	2785	N	ALA	Α	375	107.132	68.180	8.176	1.00	39.40	N
ATOM	2786	CA	ALA	A	375	106.237	67.053	8.467	1.00	39.33	C
ATOM	2787	C	ALA	Α	375	106.201	65.981	7.367	1.00	39.58	С
ATOM	2788	0	ALA	Α	375	105.988	66.289	6.194	1.00	40.42	0
ATOM	2789	CB	ALA	Α	375	104.827	67.568	8.740	1.00		С
MOTA	2790	N	PRO		376	106.397	64.701	7.742	1.00	39.51	N
ATOM	2791	CA			376	106.400	63.547	6.833	1.00		C
ATOM	2792	C			376	105.074	63.223	6.115	1.00		C
ATOM	2793	0			376	105.034	62.339	5.257	1.00	44.07	0
ATOM	2794	CB	PRO		376	106.840	62.403	7.751	1.00	38.78	C
ATOM	2795	CG	PRO			107.612	63.112	8.842	1.00	37.80	C
ATOM	2796	CD	PRO			106.665	64.236	9.111	1.00		C
ATOM	2797	N	GLY			103.992	63.916	6.465	1.00	44.20	N
ATOM	2798	CA	GLY			103.332	63.657	5.829	1.00		C
ATOM	2799	C	GLY			102.704	63.611	4.312	1.00		C
											0
ATOM	2800	O	GLY		378	103.714	64.150	3.720	1.00		N
ATOM	2801	N				101.801	62.968	3.681	1.00	48.24	C
ATOM	2802	CA			378	101.759	62.841	2.221	1.00	48.97	
ATOM	2803	C	GLU			100.987	64.015	1.627	1.00		C
ATOM	2804	0	GLU			99.905	64.345	2.096		49.36	0
ATOM	2805	CB	GLU			101.082	61.523	1.828		50.65	C
MOTA	2806	CG	GLU		378	101.750	60.266	2.409	1.00		C
ATOM	2807	CD	GLU			101.682	60.184	3.950	1.00		C
ATOM	2808	OE1	GLU			100.554	60.078	4.493		61.73	0
ATOM	2809	OE2	GLU			102.752	60.225	4.619		60.95	0
MOTA	2810	N			379	101.538	64.648	0.598	1.00	48.86	N
MOTA	2811	CA			379	100.868	65.786	-0.030	1.00		C
MOTA	2812	С			379	99.683	65.376	-0.911		49.89	C
MOTA	2813	0			379	99.745	64.394	-1.658	1.00		0
ATOM	2814	CB	THR		379	101.845	66.617	-0.903	1.00		C
ATOM	2815	OG1			379	102.921	67.089	-0.087		49.52	0
ATOM	2816	CG2	THR	A	379	101.133	67.821	-1.527	1.00		C
MOTA	2817	N	GLU	A	380	98.604	66.147	-0.814	1.00		N
MOTA	2818	CA	GLU	Α	380	97.407	65.905	-1.601	1.00		C
ATOM	2819	C	GLU	Α	380	96.897	67.228	-2.134	1.00	50.88	C
MOTA	2820	0	GLU	Α	380	97.110	68.283	-1.538	1.00		0
ATOM	2821	CB	GLU	Α	380	96.322	65.227	-0.758	1.00	51.47	C
ATOM	2822	CG	GLU	A	380	96.762	63.909	-0.168	1.00	53.35	С
ATOM	2823	CD	GLU	A	380	95.633	63.169	0.503	1.00	55.74	С
MOTA	2824	OE1	GLU	Α	380	94.962	63.764	1.373	1.00	56.99	0
MOTA	2825	OE2	GLU	A	380	95.421	61.983	0.167	1.00	57.47	0
MOTA	2826	N	ILE	A	381	96.226	67.165	-3.272	1.00	52.06	N
ATOM	2827	CA	ILE	A	381	95.697	68.364	-3.884	1.00	53.51	C
ATOM	2828	С	ILE	A	381	94.196	68.250	-4.113	1.00	53.54	С
MOTA	2829	0	ILE	A	381	93.707	67.300	-4.731	1.00	53.35	0



ATOM	2830	CB	ILE	Α	381	96.436	68.676	-5.219	1.00 54.46	C
ATOM	2831	CG1	ILE	Α	381	96.307	67.515	-6.199	1.00 54.90	C
MOTA	2832	CG2	ILE	Α	381	97.919	68.895	-4.947	1.00 55.14	C
ATOM	2833	CD1	ILE	Α	381	97.141	67.712	-7.458	1.00 55.73	C
ATOM	2834	N	TYR	A	382	93.472	69.228	-3.579	1.00 53.11	N
MOTA	2835	CA	TYR	Α	382	92.029	69.285	-3.699	1.00 52.90	C
ATOM	2836	C	TYR	Α	382	91.669	70.747	-3.928	1.00 52.45	C
ATOM	2837	0	TYR	Α	382	92.092	71.623	-3.173	1.00 51.87	0
ATOM	2838	CB	TYR	Α	382	91.379	68.732	-2.419	1.00 53.99	C
ATOM	2839	CG	TYR	Α	382	89.868	68.819	-2.388	1.00 55.14	C
ATOM	2840	CD1	TYR	Α	382	89.221	69.793	-1.623	1.00 55.14	C
ATOM	2841	CD2	TYR	Α	382	89.084	67.957	-3.161	1.00 55.70	C
ATOM	2842	CE1	TYR	Α	382	87.827	69.910	-1.629	1.00 56.22	C
ATOM	2843	CE2	TYR	Α	382	87.688	68.066	-3.177	1.00 56.34	C
ATOM	2844	CZ	TYR	A	382	87.066	69.046	-2.410	1.00 56.68	C
ATOM	2845	ОН	TYR	A	382	85.693	69.174	-2.439	1.00 57.02	0
ATOM	2846	N	GLN	Α	383	90.903	70.999	-4.987	1.00 52.24	N
ATOM	2847	CA	GLN	Α	383	90.484	72.348	-5.358	1.00 52.02	C
ATOM	2848	C	GLN	Α	383	91.687	73.210	-5.721	1.00 51.03	C
ATOM	2849	0	GLN	A	383	91.619	74.436	-5.661	1.00 50.67	0
ATOM	2850	CB	GLN	Α	383	89.707	73.018	-4.218	1.00 53.91	C
ATOM	2851	CG	GLN	Α	383	88.436	72.297	-3.794	1.00 56.88	C
ATOM	2852	CD	GLN	A	383	87.451	72.115	-4.934	1.00 59.26	C
ATOM	2853	OE1	GLN	A	383	87.746	71.441	-5.926	1.00 60.79	0
ATOM	2854	NE2	GLN	Α	383	86.270	72.718	-4.800	1.00 59.95	N
ATOM	2855	N	GLY	A	384	92.788	72.562	-6.091	1.00 50.37	N
ATOM	2856	CA	GLY	A	384	93.985	73.292	-6.468	1.00 50.11	C
ATOM	2857	С	GLY	Α	384	94.977	73.527	-5.341	1.00 49.99	C
ATOM	2858	0	GLY	Α	384	96.185	73.637	-5.586	1.00 50.26	0
ATOM	2859	N	ARG	Α	385	94.480	73.610	-4.109	1.00 49.41	N
ATOM	2860	CA	ARG	Α	385	95.350	73.837	-2.958	1.00 48.78	C
ATOM	2861	C	ARG	Α	385	96.049	72.559	-2.527	1.00 47.90	C
ATOM	2862	0	ARG	Α	385	95.536	71.456	-2.727	1.00 47.68	0
ATOM	2863	CB	ARG	Α	385	94.563	74.379	-1.761	1.00 49.61	C
MOTA	2864	CG	ARG	Α	385	93.830	75.696	-1.980	1.00 51.31	C
MOTA	2865	CD	ARG	Α	385	92.522	75.518	-2.727	1.00 52.37	C
MOTA	2866	NE	ARG	Α	385	91.851	76.801	-2.898	1.00 54.89	N
MOTA	2867	CZ	ARG	A	385	90.660	76.958	-3.467	1.00 56.27	C
MOTA	2868	NH1	ARG	Α	385	89.996	75.905	-3.925	1.00 57.58	N
MOTA	2869	NH2	ARG	A	385	90.135	78.170	-3.581	1.00 55.72	N
ATOM	2870	N	LYS	A	386	97.220	72.712	-1.920	1.00 47.18	N
MOTA	2871	CA	LYS	A	386	97.977	71.561	-1.450	1.00 46.53	C
MOTA	2872	C	LYS	A	386	97.702	71.259	0.022	1.00 45.66	C
ATOM	2873	0			386	97.602	72.166	0.857	1.00 45.07	0
MOTA	2874	CB			386	99.478	71.781	-1.642	1.00 46.65	C
ATOM	2875	CG			386	99.942	71.885	-3.082	1.00 48.06	C
ATOM	2876	CD	LYS	Α	386	101.451	72.061	-3.101	1.00 50.09	С

MOTA

2923

CB

### **ATOM** 2877 CE LYS A 386 102.017 -4.505 1.00 51.55 72.118 C **ATOM** 2878 LYS A 386 103.505 72.226 NZ1.00 52.03 -4.438 И **ATOM** TYR A 387 97.580 2879 N 1.00 45.21 69.971 0.326 N **ATOM** 2880 TYR A 387 97.335 CA 69.515 1.689 1.00 45.09 C **ATOM** 2881 **TYR A 387** 98.420 C 68.561 2.155 1.00 44.05 TYR A 387 **ATOM** 2882 99.270 68.119 1.380 1.00 44.31 **ATOM TYR A 387** 95.978 2883 68.807 CB1.00 45.22 1.787 C **TYR A 387 ATOM** 2884 CG 94.799 69.725 1.580 1.00 45.09 CD1 TYR A 387 ATOM 2885 94.537 0.332 1.00 45.10 70.291 CD2 TYR A 387 **ATOM** 2886 93.981 70.083 2.652 1.00 45.00 C **ATOM** 2887 **CE1 TYR A 387** 93.492 0.160 71.194 C 1.00 45.13 **ATOM** 2888 TYR A 387 CE2 92.941 70.979 2.491 C 1.00 45.41 **TYR A 387** ATOM 2889 CZ92.707 1.00 45.60 71.531 1.246 C **ATOM** 2890 TYR A 387 OH 91.701 72.444 1.00 47.56 1.104 0 **MOTA** LYS A 388 2891 N 68.248 3.438 98.381 1.00 43.76 $\mathbf{N}$ LYS A 388 **ATOM** 2892 CA 67.326 4.034 99.328 1.00 43.49 C **ATOM** 2893 LYS A 388 98.501 4.865 66.389 1.00 41.84 C **ATOM** 2894 LYS A 388 97.731 66.834 5.703 1.00 40.70 0 2895 LYS A 388 **ATOM** CB 100.317 1.00 45.64 68.067 4.932 C 2896 LYS A 388 MOTA CG 101.504 68.715 4.213 C 1.00 48.31 **ATOM** 2897 LYS A 388 102.626 CD3.892 67.706 C 1.00 47.96 **ATOM** 2898 LYS A 388 CE 103.901 68.437 3.466 1.00 47.56 C **ATOM** 2899 LYS A 388 105.097 67.563 3.426 NZ1.00 46.67 N THR A 389 **ATOM** 2900 65.092 4.617 N 98.647 1,00 42.01 N **ATOM** 2901 THR A 389 97.902 64.098 5.370 CA C 1.00 41.89 MOTA 2902 THR A 389 98.290 6.833 1.00 42.40 C C 64.224 **ATOM** THR A 389 99.418 2903 0 64.598 1.00 42.02 7.172 THR A 389 **ATOM** 2904 4.909 98.218 62.669 CB 1.00 41.09 C **ATOM** 2905 OG1 THR A 389 62.427 5.042 99.623 1.00 41.52 CG2 THR A 389 97.807 **ATOM** 2906 62.474 3.461 1.00 41.23 MOTA 2907 TYR A 390 97.330 63.930 7.693 1.00 42.51 MOTA 2908 CA TYR A 390 97.533 63.980 9.125 1.00 42.75 C 2909 TYR A 390 62.882 9.670 MOTA C 96.624 1.00 43.43 ATOM TYR A 390 0 2910 95.406 62.969 9.551 1.00 44.28 ATOM TYR A 390 2911 СB C 97.098 65.332 9.679 1.00 41.44 2912 MOTA CG TYR A 390 97.362 65.467 C 11.159 1.00 40.95 ATOM 96.609 C 2913 CD1 TYR A 390 66.343 11.944 1.00 41.25 CD2 TYR A 390 64.729 11.777 C ATOM 2914 98.373 1.00 39.46 C ATOM 96.852 66.478 2915 CE1 TYR A 390 13.312 1.00 40.17 98.623 MOTA 2916 CE2 TYR A 390 64.856 13.136 1.00 40.26 C TYR A 390 MOTA 2917 CZ97.857 65.732 13.896 1.00 39.92 **ATOM** 2918 TYR A 390 98.095 15.238 $\mathsf{OH}$ 65.857 1.00 40.43 N **ATOM** 2919 ARG A 391 97.206 61.846 10.258 1.00 44.01 N ARG A 391 10.772 **ATOM** 2920 CA 96.399 60.750 1.00 44.87 ARG A 391 C ATOM 96.778 2921 60.387 12.202 1.00 45.41 2922 MOTA ARG A 391 97.954 12.580 1.00 45.88 60.450 ARG A 391

96.589 59.529

9.883

1.00 44.93

### 1.00 46.56 **MOTA** 2924 CG ARG A 391 97.990 58.967 9.986 C ARG A 391 $\mathbf{C}$ 98.242 1.00 47.95 MOTA 2925 CD57.872 8.981 ARG A 391 9.200 N ATOM 2926 NE 99.535 57.232 1.00 48.70 ARG A 391 CZC ATOM 2927 100.072 56.357 8.363 1.00 49.29 2928 56.024 7.252 1.00 50.30 NH1 ARG A 391 99.425 N MOTA NH2 ARG A 391 8.638 2929 101.245 55.805 1.00 49.69 MOTA N **GLY A 392** 95.782 **ATOM** 2930 60.009 12.999 N 1.00 45.56 N **GLY A 392** 59.618 C **ATOM** 2931 CA 96.062 14.366 1.00 45.33 **GLY A 392** 96.803 58.291 **ATOM** 2932 C 14.343 1.00 45.81 13.451 **ATOM** 2933 **GLY A 392** 96.578 57.467 1.00 45.18 0 MSE A 393 97.696 1.00 46.09 58.074 15.304 $\mathbf{N}$ HETATM 2934 N HETATM 2935 MSE A 393 98.437 56.818 15.344 1.00 46.91 C CA MSE A 393 97.514 C **HETATM 2936** 55.638 15.668 1.00 45.83 HETATM 2937 MSE A 393 97.905 15.553 54.474 0 1.00 45.29 HETATM 2938 MSE A 393 56.925 16.351 1.00 48.46 C CB 99.587 MSE A 393 **HETATM 2939** 100.671 57.905 15.906 1.00 51.44 C CG SE HETATM 2940 SE MSE A 393 101.600 57.363 14.403 1.00 55.48 MSE A 393 **HETATM 2941** 102.496 55.927 15.071 1.00 55.49 C CE **ATOM GLY A 394** N 2942 N 96.280 55.954 16.049 1.00 44.70 54.926 **GLY A 394** 95.307 16.364 1.00 44.19 C **ATOM** 2943 CA **GLY A 394** 94.355 54.686 15.206 1.00 44.14 C **MOTA** 2944 C 93.416 2945 **GLY A 394** 53.896 0 MOTA 0 15.311 1.00 43.61 **SER A 395 ATOM** N 2946 N 94.591 55.369 14.093 1.00 44.85 SER A 395 93.740 55.204 1.00 45.69 C **ATOM** 2947 12.921 CA C **ATOM** 2948 SER A 395 C 94.061 53.866 12.265 1.00 46.80 **ATOM** SER A 395 53.175 12.663 1.00 47.02 0 2949 0 95.002 1.00 45.01 SER A 395 93.968 56.340 C ATOM 2950 CB 11.916 **SER A 395** 56.305 95.278 11.372 1.00 44.71 ATOM 2951 OG ILE A 396 93.276 11.261 1.00 47.68 **ATOM** 2952 53.501 N N C ILE A 396 93.488 52.245 10.554 1.00 49.11 ATOM 2953 CA 52.246 2954 ILE A 396 94.830 9.797 1.00 49.53 ATOM 0 ILE A 396 95.705 51.418 10.070 1.00 48.83 MOTA 2955 ILE A 396 C ATOM 92.320 51.974 9.558 1.00 49.53 2956 CB C 90.990 2957 CG1 ILE A 396 51.908 10.315 1.00 49.28 MOTA **ATOM** 2958 92.554 50.670 8.812 1.00 49.14 C CG2 ILE A 396 50.820 C **ATOM** 90.932 11.363 1.00 49.48 2959 CD1 ILE A 396 N 53.183 8.863 **ATOM** ALA A 397 94.988 1.00 50.24 2960 C 1.00 51.94 CA ALA A 397 **ATOM** 53.291 8.062 2961 96.206 1.00 53.51 C ATOM 2962 8.903 C ALA A 397 97.475 53.255 0 52.533 ATOM 2963 ALA A 397 98.414 8.576 1.00 53.35 C ALA A 397 7.227 1.00 50.87 **ATOM** 96.177 54.567 2964 CB N 54.037 ALA A 398 97.507 9.979 1.00 55.56 ATOM 2965 N C 10.851 2966 ALA A 398 98.677 54.074 1.00 57.66 ATOM CA C **ATOM** 98.878 52.719 11.529 1.00 59.35 2967 ALA A 398 0 1.00 59.25 2968 ALA A 398 99.988 52.181 11.541 ATOM C 55.175 11.900 1.00 56.88 ALA A 398 98.525 ATOM 2969 CBN 97.803 52.162 12.084 1.00 61.57 HETATM 2970 MSE A 399

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HETATM	2971	CA	MSE	A	399	97.883	50.868	12.753	1.00	63.53	C
HETATM	2972	C	MSE	A	399	98.165	49.736	11.778	1.00	64.22	C
${\tt HETATM}$	2973	0	MSE	Α	399	98.284	48.586	12.181	1.00	64.45	0
HETATM	2974	CB	MSE	A	399	96.589	50.561	13.510	1.00	64.53	C
HETATM	2975	CG	MSE	A	399	96.335	51.428	14.726	1.00	66.17	C
HETATM	2976	SE	MSE	Α	399	94.927	50.801	15.688	1.00	70.47	SE
HETATM	2977	CE	MSE	A	399	93.579	50.893	14.454	1.00	68.03	C
MOTA	2978	N	LYS	A	400	98.275	50.067	10.498	1.00	65.70	N
MOTA	2979	CA	LYS	Α	400	98.535	49.076	9.461	1.00	67.25	C
MOTA	2980	C	LYS	Α	400	99.753	48.195	9.743	1.00	68.24	C
MOTA	2981	0	LYS	Α	400	100.162	48.015	10.888	1.00	68.63	0
MOTA	2982	CB	LYS	Α	400	98.703	49.771	8.118	1.00	67.70	C
MOTA	2983	N	LYS	Α	401	100.324	47.646	8.677	1.00	69.15	N
MOTA	2984	CA	LYS	Α	401	101.487	46.771	8.773	1.00	70.20	C
MOTA	2985	C	LYS	A	401	102.588	47.357	9.662	1.00	71.05	C
MOTA	2986	0	LYS	A	401	102.394	48.471	10.197	1.00	71.27	0
MOTA	2987	CB	LYS	Α	401	102.035	46.483	7.365	1.00	69.48	C
ATOM	2988	N	ASN	A	416	92.074	47.487	5.186	1.00	72.24	N
MOTA	2989	CA	ASN	Α	416	91.155	46.876	6.194	1.00	72.64	C
MOTA	2990	C	ASN	A	416	91.165	47.666	7.501	1.00	72.60	C
MOTA	2991	0	ASN	Α	416	92.209	48.176	7.915	1.00	73.31	0
ATOM	2992	CB	ASN	Α	416	91.563	45.422	6.465	1.00	72.21	C
ATOM	2993	N	LYS	A	417	90.004	47.744	8.152	1.00	71.58	N
MOTA	2994	CA	LYS	Α	417	89.858	48.463	9.419	1.00	70.32	C
ATOM	2995	C	LYS	Α	417	89.985	47.484	10.580	1.00	69.53	C
MOTA	2996	0	LYS	A	417	89.416	46.386	10.515	1.00	70.17	0
ATOM	2997	CB	LYS	Α	417	88.499	49.146	9.472	1.00	70.50	C
ATOM	2998	N	LEU	A	418	90.712	47.870	11.642	1.00	67.91	N
ATOM	2999	CA	LEU	Α	418	90.903	46.996	12.826	1.00	64.90	Ċ
MOTA	3000	C	LEU	Α	418	90.637	47.647	14.211	1.00	62.37	C
MOTA	3001	0	LEU	Α	418	91.478	47.559	15.111	1.00	62.60	0
ATOM	3002	CB	LEU	A	418	92.286	46.414	12.785	1.00	64.79	C
ATOM	3003	N	VAL	Α	419	89.455	48.260	14.362	1.00	59.18	N
ATOM	3004	CA	VAL	A	419	88.983	48.973	15.576	1.00	56.04	C
ATOM	3005	C	VAL	Α	419	89.820	50.195	15.995	1.00	53.39	C
ATOM	3006	0	VAL	Α	419	90.533	50.180	17.005	1.00	51.68	0
MOTA	3007	CB	VAL	A	419	88.825	48.027	16.790	1.00	56.09	C
MOTA	3008	CG1	VAL	A	419	88.330	48.818	18.006	1.00	55.27	C
ATOM	3009	CG2	VAL	A	419	87.823	46.925	16.456	1.00	56.32	С
ATOM	3010	N	PRO	A	420	89.728	51.280	15.207	1.00	51.10	N
MOTA	3011	CA	PRO	A	420	90.430	52.548	15.402	1.00	49.87	С
MOTA	3012	C	PRO	A	420	89.937	53.428	16.559		48.67	С
MOTA	3013	0	PRO	A	420	88.758	53.419	16.924		47.55	0
MOTA	3014	CB	PRO	A	420	90.238	53.225	14.052		49.97	C
MOTA	3015	CG	PRO	A	420	88.824	52.828	13.734		49.16	C
MOTA	3016	CD	PRO	A	420	88.939	51.338	13.961		49.81	C
MOTA	3017	N	GLU	A	421	90.871	54.192	17.116	1.00	48.34	N

### **GLU A 421** 1.00 47.97 CA 90.610 55.123 18.208 C ATOM 3018 3019 90.954 56.506 17.668 1.00 46.84 C MOTA C **GLU A 421** 91.005 **ATOM** 3020 0 0 GLU A 421 57.481 18.409 1.00 47.51 19.396 **GLU A 421** 91.517 54.809 MOTA 3021 CB 1.00 49.79 C 91.207 3022 C CG GLU A 421 53.512 20.149 1.00 53.19 ATOM C **ATOM** 3023 CD **GLU A 421** 90.329 53.739 21.372 1.00 53.99 54.563 1.00 55.23 3024 OE1 GLU A 421 90.730 22.228 0 ATOM OE2 GLU A 421 89.260 MOTA 0 3025 53.095 21.484 1.00 53.71 1.00 45.35 ATOM 3026 **GLY A 422** 91.200 56.576 N N 16.366 1.00 44.97 3027 **GLY A 422** 91.541 57.834 MOTA 15.736 C CA **GLY A 422** 91.214 3028 57.813 14.253 1.00 44.86 C MOTA C **GLY A 422** 13.684 1.00 44.42 90.918 56.760 **MOTA** 3029 0 0 3030 ILE A 423 91.279 58.974 13.611 1.00 44.45 N MOTA N 59.045 ILE A 423 90.954 12.196 1.00 43.46 C MOTA 3031 CA C 3032 ILE A 423 92.129 59.498 11.321 1.00 42.96 MOTA C 3033 ILE A 423 93.178 0 ATOM 59.916 11.821 1.00 41.95 0 ILE A 423 89.703 59.963 11.970 1.00 43.06 C 3034 CB MOTA 3035 61.412 C CG1 ILE A 423 89.978 12.403 1.00 42.83 MOTA C CG2 ILE A 423 88.534 59.439 12.788 **ATOM** 3036 1.00 41.55 90.868 C **ATOM** 1.00 43.42 3037 CD1 ILE A 423 62.208 11.461 91.934 59.387 1.00 42.19 N MOTA 3038 Ν GLU A 424 10.012 **GLU A 424** 92.931 59.775 C **ATOM** 9.024 1.00 42.21 3039 CA **GLU A 424** 92.422 61.047 C **ATOM** 3040 8.363 1.00 41.48 C 3041 **GLU A 424** 91.266 7.945 1.00 41.26 0 61.110 MOTA 0 93.071 7.963 1.00 43.26 C MOTA 3042 CB GLU A 424 58.680 **GLU A 424** 93.518 C MOTA 57.325 8.492 1.00 45.35 3043 CG 56.225 1.00 46.93 C 93.357 7.453 MOTA 3044 CD GLU A 424 56.423 OE1 GLU A 424 93.818 6.305 1.00 48.20 0 3045 MOTA 92.780 OE2 GLU A 424 7.780 1.00 46.63 0 **ATOM** 55.162 3046 93.276 62.060 8.270 N N **GLY A 425** 1.00 41.08 ATOM 3047 92.857 63.303 1.00 40.46 **ATOM** GLY A 425 7.643 3048 ÇA C GLY A 425 93.976 64.030 6.920 1.00 40.17 ATOM 3049 C 0 6.609 1.00 39.82 GLY A 425 95.017 63.446 MOTA 3050 O 65.309 ARG A 426 6.643 1.00 39.48 N MOTA 3051 93.757 N 66.117 5.967 1.00 39.94 C ARG A 426 94.753 ATOM 3052 CA C 94.571 6.375 3053 67.562 1.00 40.26 MOTA C ARG A 426 Q ARG A 426 67.963 6.792 1.00 41.45 **ATOM** 93.486 3054 0 C 65.989 4.452 1.00 39.82 3055 94.608 MOTA CB ARG A 426 1.00 41.77 C 3.882 CG ARG A 426 93.320 66.554 **ATOM** 3056 2.405 C 93.202 66.182 1.00 43.77 3057 CD ARG A 426 ATOM 1.00 45.14 N 3058 NE ARG A 426 92.026 66.747 1.751 MOTA C 0.506 1.00 46.18 91.654 66.455 CZ ARG A 426 **ATOM** 3059 92.366 65.604 N -0.218 1.00 46.99 NH1 ARG A 426 3060 MOTA N -0.021 1.00 47.23 90.577 67.021 3061 NH2 ARG A 426 ATOM N 6.267 1.00 40.61 VAL A 427 95.640 68.342 ATOM 3062 N C 95.592 6.613 1.00 40.23 69.759 MOTA CA VAL A 427 3063 C 5.548 3064 C VAL A 427 96.338 70.557 1.00 40.03 MOTA

ATOM	3065	0	VAL	A	427	97.235	70.033	4.885	1.00	39.67	0
ATOM	3066	CB	VAL	Α	427	96.225	70.029	8.001	1.00	39.28	C
MOTA	3067	CG1	VAL	Α	427	95.457	69.280	9.079	1.00	37.97	C
ATOM	3068	CG2	VAL	A	427	97.681	69.609	7.993	1.00	39.20	C
ATOM	3069	N	ALA	A	428	95.945	71.815	5.381	1.00	40.23	N
MOTA	3070	CA	ALA	A	428	96.562	72.700	4.406	1.00	41.04	C
MOTA	3071	C	ALA	A	428	98.079	72.829	4.595	1.00	42.53	C
MOTA	3072	0	ALA			98.578	72.780	5.727		42.82	0
ATOM	3073	CB	ALA			95.918	74.074	4.496	1.00		C
ATOM	3074	Ŋ	TYR			98.801	72.975	3.480		42.74	N
MOTA	3075	CA	TYR			100.251	73.168	3.491		43.20	C
ATOM	3076	C	TYR			100.546	74.452	4.284		42.09	C
MOTA	3077	O	TYR			99.986	75.509	3.984		41.87	0
ATOM	3078	CB	TYR			100.757	73.332	2.061		44.90	C
ATOM ATOM	3079 3080	CG CD1	TYR TYR			102.211 103.228	73.730 72.842	1.972 2.329	1.00	48.81 50.14	C
ATOM	3080	CD2	TYR			103.228	75.005	1.541		49.88	C
ATOM	3082	CE1	TYR			104.575	73.003	2.255		50.62	C
ATOM	3083	CE2	TYR			103.915	75.388	1.463	1.00		C
ATOM	3084	CZ	TYR			104.908	74.487	1.819	1.00		C
ATOM	3085	OH	TYR			106.226	74.863	1.718	1.00		0
ATOM	3086	N	LYS	A	430	101.429	74.372	5.276	1.00	40.05	N
ATOM	3087	CA	LYS	A	430	101.727	75.541	6.102	1.00	38.56	C
ATOM	3088	C	LYS	A	430	103.086	76.198	5.889	1.00	37.11	C
MOTA	3089	0	LYS	A	430	103.360	77.238	6.486	1.00	37.51	0
MOTA	3090	CB	LYS	A	430	101.610	75.178	7.589	1.00	38.70	C
ATOM	3091	CG	LYS	A	430	100.294	74.563	7.981	1.00	38.40	C
MOTA	3092	CD	LYS	Α	430	100.284	74.152	9.434	1.00		C
MOTA	3093	CE	LYS			98.997	73.409	9.738		39.27	C
ATOM	3094	NZ	LYS			98.927	72.970	11.149		40.88	N
ATOM	3095	N	GLY			103.942	75.609	5.063	1.00		N
ATOM	3096	CA	GLY			105.256	76.199	4.867	1.00		C
ATOM	3097	C	GLY			106.199	75.798	5.991 6.710		31.20 30.24	C O
ATOM ATOM	3098 3099	O N	GLY ALA			105.927 107.296	74.847 76.528	6.157	1.00		N
ATOM	3100	CA	ALA			108.290	76.214	7.189	1.00		C
ATOM	3101	C	ALA			107.759	76.197	8.618		29.06	C
ATOM	3102	0	ALA			106.981	77.057	9.018	1.00		0
ATOM	3103	CB	ALA			109.462	77.188	7.097	1.00		C
ATOM	3104	N	ALA			108.206	75.216	9.390	1.00	27.87	N
ATOM	3105	CA	ALA	Α	433	107.798	75.077	10.777	1.00	27.30	C
ATOM	3106	С	ALA	A	433	108.341	76.225	11.624	1.00	27.35	C
MOTA	3107	0	ALA	A	433	107.727	76.611	12.620	1.00	27.15	0
MOTA	3108	CB	ALA	Α	433	108.289	73.739	11.333	1.00	28.08	C
ATOM	3109	N			434	109.493	76.766	11.238		26.71	Ŋ
MOTA	3110	CA	SER			110.085	77.875	11.983		27.06	C
MOTA	3111	C	SER	Α	434	109.131	79.070	12.036	1.00	26.79	C



109.120 79.821 13.014 1.00 25.57 0 SER A 434 ATOM 3112 0 78.306 11.357 C 1.00 27.17 **SER A 434** 111.419 **ATOM** 3113 CB **ATOM** 3114 **SER A 434** 111.252 78.778 10.031 1.00 28.63 0 OG N **ATOM ASP A 435** 108.333 79.243 10.986 1.00 27.24 3115 N C **ATOM** 3116 **ASP A 435** 80.333 10.928 1.00 28.29 CA 107.363 **ASP A 435** C 1.00 27.92 C 106.188 80.060 11.868 ATOM 3117 1.00 28.07 3118 **ASP A 435** 105.714 80.959 12.565 ATOM 0 **ASP A 435** 80.517 9.491 C 106.876 1.00 31.68 MOTA 3119 CB**ASP A 435** 1.00 35.77 C **ATOM** 3120 CG 107.949 81.120 8.579 OD1 ASP A 435 107.761 81.087 0 **ATOM** 3121 7.341 1.00 38.23 1.00 36.12 **ATOM** OD2 ASP A 435 108.968 81.643 9.096 0 3122 ILE A 436 11.886 N 3123 78.815 1.00 26.54 **ATOM** N 105.726 1.00 25.99 C 3124 ILE A 436 104.637 78.398 12.763 CA MOTA ILE A 436 78.679 C 1.00 24.91 **ATOM** 3125 105.085 14.199 C 79.320 1.00 23.37 3126 ILE A 436 14.977 0 104.388 MOTA О ILE A 436 76.867 12.617 C 1.00 26.46 MOTA 104.357 3127 CBCG1 ILE A 436 11.166 1.00 28.02 C 3128 104.016 76.524 ATOM CG2 ILE A 436 103.236 76.437 C 3129 13.536 1.00 26.46 ATOM C 1.00 30.24 3130 CD1 ILE A 436 102.860 77.292 10.604 ATOM **VAL A 437** N 14.539 1.00 25.92 **ATOM** 3131 N 106.268 78.191 **VAL A 437** C 3132 106.804 78.383 15.877 1.00 27.27 ATOM CA C 79.853 VAL A 437 16.244 1.00 28.27 ATOM 106.902 3133 C 0 3134 VAL A 437 106.560 80.238 17.367 1.00 29.71 0 MOTA 1.00 26.18 **VAL A 437** 77.747 16.022 C 108.200 MOTA 3135 CB 108.770 17.395 1.00 25.87 C 3136 CG1 VAL A 437 78.055 ATOM 1.00 26.99 C 76.244 15.821 CG2 VAL A 437 108.105 MOTA 3137 PHE A 438 107.367 80.678 15.307 1.00 29.44 N N ATOM 3138 107.513 1.00 29.34 C PHE A 438 82.104 15.564 3139 **ATOM** CA 1.00 29.77 82.759 15.872 C PHE A 438 **MOTA** C 106.182 3140 83.647 106.110 16.721 1.00 29.60 0 PHE A 438 3141 ATOM 108.183 CB82.799 1.00 30.41 MOTA 3142 PHE A 438 14.382 C 14.589 1.00 32.43 CG PHE A 438 108.391 84.262 MOTA 3143 C 1.00 34.66 CD1 PHE A 438 107.342 85.160 14.424 ATOM 3144 C 109.625 15.012 1.00 34.35 MOTA CD2 PHE A 438 84.742 3145 C 107.514 1.00 35.25 CE1 PHE A 438 86.522 14.678 ATOM 3146 15.273 C 1.00 35.59 109.816 86.100 CE2 PHE A 438 ATOM 3147 C PHE A 438 108.752 86.994 15.105 1.00 36.08 ATOM 3148 CZ1.00 31.03 105.125 15.189 N GLN A 439 82.325 N ATOM 3149 C 1.00 31.99 3150 103.801 82.885 15.440 ATOM CAGLN A 439 C 103.280 82.473 1.00 32.50 16.803 GLN A 439 ATOM 3151 C 0 102.669 GLN A 439 83.282 17.502 1.00 33.67 MOTA 3152 0 82.454 C 14.369 1.00 33.33 **ATOM** 3153 GLN A 439 102.801 CB C 83.175 1.00 36.37 GLN A 439 102.962 13.052 MOTA 3154 CG C 13.204 1.00 38.29 GLN A 439 102.863 84.685 ATOM 3155 CD0 1.00 39.62 MOTA 3156 OE1 GLN A 439 101.882 85.212 13.746 1.00 39.30 85.393 12.720 NE2 GLN A 439 103.880 ATOM 3157 1.00 31.91 N 81.225 17.199 N MSE A 440 103.519 HETATM 3158

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HETATM	3159	CA	MSE	A	440	103.036	80.797	18.498	1.00	32.48	C
HETATM	3160	С	MSE	A	440	103.802	81.394	19.682	1.00	30.42	C
HETATM	3161	0	MSE	A	440	103.199	81.680	20.718	1.00	29.98	0
HETATM	3162	CB	MSE	Α	440	102.958	79.261	18.577	1.00	36.81	C
HETATM	3163	CG	MSE	A	440	104.200	78.483	18.205	1.00	44.35	C
HETATM	3164	SE	MSE	A	440	103.911	76.661	18.242	1.00	53.13	SE
HETATM	3165	CE	MSE	Α	440	102.652	76.462	16.958	1.00	51.30	С
MOTA	3166	N	LEU			105.107	81.620	19.543		27.53	N
MOTA	3167	CA	LEU		441	105.867	82.206	20.648	1.00	25.82	C
MOTA	3168	C	LEU			105.407	83.644	20.911		25.88	C
ATOM	3169	0	LEU		441	105.348	84.092	22.053	1.00		0
MOTA	3170	CB	LEU		441	107.370	82.194	20.353	1.00		C
ATOM	3171	CG	LEU		441	108.062	80.839	20.155	1.00		C
ATOM	3172	CD1			441	109.552	81.080	19.937		25.09	C
ATOM	3173	CD2			441	107.849	79.940	21.362		25.75	C
ATOM	3174	N	GLY			105.076	84.366	19.850	1.00		N C
ATOM	3175	CA	GLY			104.621	85.731	20.019	1.00		C
ATOM	3176	C O	GLY GLY			103.358	85.780 86.656	21.693	1.00		0
ATOM ATOM	3177 3178	N	GLY			103.200	84.834	20.605	1.00		N
ATOM	3179	CA	GLY			102.430	84.775	21.347		24.02	C
MOTA	3180	C	GLY			101.449	84.354	22.781		24.08	C
ATOM	3181	0	GLY			100.773	84.822	23.696		23.64	0
ATOM	3182	N	ILE			102.419	83.465	22.970	1.00		N
ATOM	3183	CA			444	102.780	82.979	24.300	1.00		С
ATOM	3184	C	ILE	A	444	103.376	84.135	25.106	1.00	24.11	C
MOTA	3185	0	ILE	Α	444	103.005	84.346	26.261	1.00	23.54	0
MOTA	3186	CB	ILE	Α	444	103.798	81.801	24.209	1.00	23.72	C
ATOM	3187	CG1	ILE	A	444	103.152	80.622	23.466	1.00	22.82	C
MOTA	3188	CG2	ILE	Α	444	104.241	81.360	25.612	1.00	22.84	С
MOTA	3189	CD1	ILE	A	444	104.110	79.479	23.146	1.00	22.36	C
ATOM	3190	N	ARG			104.289	84.890	24.500		23.72	N
MOTA	3191	CA	ARG			104.883	86.030	25.195		24.00	C
ATOM	3192	C	ARG			103.837	87.106	25.514	1.00		C
MOTA	3193	0	ARG			103.891	87.721	26.576	1.00		0
ATOM	3194	CB	ARG			106.026	86.625	24.372		22.89	C
ATOM	3195	CG	ARG			107.275	85.752	24.332		24.88	C
ATOM	3196	CD	ARG			108.356 109.592	86.373 85.596	23.456 23.462	1.00	26.53 27.53	N
ATOM ATOM	3197 3198	NE CZ	ARG ARG			110.470	85.579	24.464	1.00		C
ATOM	3199	NH1	ARG			110.262	86.303	25.561	1.00		N
ATOM	3200	NH2			445	111.559	84.823	24.372	1.00		N
ATOM	3201	N			446	102.882	87.333	24.615	1.00		N
ATOM	3202	CA			446	101.835		24.878		24.75	C
ATOM	3203	C			446	100.953	87.831	26.004		23.48	C
ATOM	3204	0			446	100.605	88.585	26.902		24.41	0
ATOM	3205	CB			446	100.953	88.558	23.656	1.00	25.42	C

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ATOM	3206	OG	SER	A	446	101.715	89.056	22.574	1.00 3	4.14	0
ATOM	3207	N	$\mathtt{GLY}$	A	447	100.587	86.559	25.946	1.00 2	2.70	N
ATOM	3208	CA	GLY	A	447	99.740	85.996	26.974	1.00 2	3.15	C
ATOM	3209	С	GLY	A	447	100.366	86.119	28.338	1.00 2	3.47	C
MOTA	3210	0	GLY	Α	447	99.707	86.531	29.288	1.00 2	4.01	0
HETATM	3211	N	MSE	Α	448	101.641	85.754	28.435	1.00 2	4.62	N
HETATM	3212	CA	MSE	Α	448	102.369	85.834	29.693	1.00 2	5.24	C
HETATM	3213	C	MSE	A	448	102.566	87.302	30.112	1.00 2	4.99	C
HETATM	3214	0	MSE	A	448	102.632	87.613	31.300	1.00 2	5.75	0
HETATM	3215	CB	MSE	A	448	103.704	85.090	29.569	1.00 2	7.58	C
HETATM	3216	CG	MSE	Α	448	103.522	83.600	29.220	1.00 3	1.58	C
HETATM	3217	SE	MSE	Α	448	105.040	82.559	29.094	1.00 3	7.58	SE
HETATM	3218	CE	MSE	Α	448	105.621	82.613	30.821	1.00 3	4.91	C
ATOM	3219	N	GLY	Α	449	102.639	88.208	29.145	1.00 2	4.08	N
MOTA	3220	CA	GLY	A	449	102.770	89.613	29.487	1.00 2	4.66	С
MOTA	3221	C	GLY	Α	449	101.497	90.132	30.154	1.00 2	5.65	C
MOTA	3222	0	GLY	Α	449	101.550	90.876	31.131	1.00 2	5.66	0
MOTA	3223	N	TYR	A	450	100.341	89.739	29.626	1.00 2	5.60	N
MOTA	3224	CA	TYR	Α	450	99.059	90.166	30.178	1.00 2	5.77	С
MOTA	3225	C	TYR	A	450	98.831	89.753	31.623	1.00 2	4.83	C
MOTA	3226	0	TYR	Α	450	98.211	90.485	32.375		5.34	0
ATOM	3227	CB	TYR	Α	450	97.899	89.622	29.344	_	6.05	С
MOTA	3228	CG	TYR	Α	450	97.590	90.402	28.096		7.41	C
MOTA	3229	CD1	TYR	Α	450	97.111	91.706	28.167		8.14	* <b>C</b>
MOTA	3230	CD2	TYR	Α	450	97.749	89.826	26.839		8.61	С
ATOM	3231	CE1	TYR	Α	450	96.793	92.419	27.007		9.06	С
MOTA	3232	CE2	TYR	A	450	97.436	90.530	25.678		9.51	C
ATOM	3233	CZ	TYR	Α	450	96.959	91.822	25.770		29.28	C
ATOM	3234	OH	TYR	Α	450	96.640	92.501	24.618	1.00 3		0
MOTA	3235	N	VAL	A	451	99.305	88.576	32.010		4.89	N
MOTA	3236	CA	VAL	Α	451	99.098	88.121	33.376		5.62	C
MOTA	3237	С			451	100.316	88.418	34.251		26.97	C
MOTA	3238	0			451	100.343	88.080	35.434		27.47	0
MOTA	3239	CB			451	98.758	86.605	33.417		24.80	C
ATOM	3240	CG1			451	97.465	86.346	32.657		22.49	C
ATOM	3241	CG2			451	99.884	85.787	32.822	1.00 2		C
ATOM	3242	N			452	101.318	89.061	33.655		28.11	N
MOTA	3243	CA			452	102.518	89.425	34.384		28.84	C
ATOM	3244	С			452	103.396	88.273	34.816		30.29	C
ATOM	3245	0			452	104.016	88.325	35.873		30.53	0
MOTA	3246	N			453	103.461	87.231	33.998		30.98	N
ATOM	3247	CA			453	104.278	86.065	34.312		31.22	C
ATOM	3248	C			453	105.623	86.177	33.612	1.00 3		C
ATOM	3249	0			453	105.694	86.079	32.386	1.00 3		0
ATOM	3250				453	103.567	84.800	33.862	1.00 3		C
ATOM	3251	N			454	106.686	86.380	34.387	1.00 3		N C
ATOM	3252	CA	GLY	A	454	108.015	86.497	33.806	1.00 3	50.70	C



MOTA	3253	C	GLY	A	454	108.595	85.166	33.359	1.00	30.23	C
MOTA	3254	0	GLY	A	454	109.509	85.117	32.534	1.00	29.71	0
MOTA	3255	N	ASP	Α	455	108.076	84.086	33.930	1.00	30.07	N
MOTA	3256	CA	ASP	Α	455	108.504	82.742	33.583	1.00	30.58	C
ATOM	3257	C	ASP	Α	455	107.326	81.825	33.830	1.00	30.31	C
MOTA	3258	0	ASP	Α	455	106.353	82.228	34.463	1.00	30.69	0
MOTA	3259	CB	ASP	A	455	109.701	82.290	34.426	1.00	32.56	C
MOTA	3260	CG	ASP	Α	455	109.415	82.301	35.919	1.00	34.66	C
MOTA	3261	OD1	ASP	A	455	108.311	81.896	36.331	1.00	36.16	0
ATOM	3262	OD2	ASP	A	455	110.313	82.688	36.694	1.00	38.54	0
MOTA	3263	N	ILE	Α	456	107.412	80.593	33.343	1.00	29.95	N
MOTA	3264	CA	ILE	Α	456	106.326	79.634	33.502	1.00	30.50	C
ATOM	3265	C	ILE	А	456	105.976	79.367	34.957	1.00	31.87	C
MOTA	3266	0	ILE	Α	456	104.820	79.103	35.274	1.00	32.52	0
MOTA	3267	CB	ILE	Α	456	106.657	78.302	32.791	1.00	29.40	C
MOTA	3268	CG1	ILE	A	456	106.775	78.555	31.287	1.00	29.89	C
MOTA	3269	CG2	ILE	Α	456	105.589	77.259	33.082	1.00	28.25	C
ATOM	3270	CD1	ILE	Α	456	107.179	77.342	30.477	1.00	31.58	C
MOTA	3271	N	GLN	A	457	106.964	79.445	35.844	1.00	32.73	N
MOTA	3272	CA	GLN	Α	457	106.718	79.207	37.264	1.00	33.99	C
MOTA	3273	C	GLN	A	457	105.672	80.193	37.801	1.00	33.89	С
MOTA	3274	0	GLN	A	457	104.718	79.800	38.477		33.85	0
ATOM	3275	CB	GLN	A	457	108.021	79.349	38.049	1.00		C
MOTA	3276	CG	GLN	Α	457	107.883	79.030	39.528	1.00		C
MOTA	3277	CD	GLN			109.173	79.257	40.300		44.64	C
ATOM	3278	OE1	GLN			110.235	78.745	39.930	1.00		0
MOTA	3279	NE2	GLN			109.084	80.019	41.386	1.00		N
MOTA	3280	N	GLU			105.860	81.475	37.494	1.00		N
ATOM	3281	CA	GLU			104.937	82.523	37.925	1.00		C
ATOM	3282	C			458	103.574	82.270	37.315	1.00		C
ATOM	3283	0	GLU			102.542	82.607	37.889	1.00		0
MOTA	3284	CB	GLU			105.436	83.899	37.478	1.00		C
ATOM	3285	CG	GLU			106.617	84.441	38.255	1.00		C
ATOM	3286	CD	GLU			107.151	85.736	37.661	1.00		0
ATOM	3287	OE1			458	106.336	86.630	37.337		39.29	0
ATOM	3288	OE2	GLU			108.390	85.867	37.532	1.00		N
ATOM	3289	N			459	103.587	81.673	36.137	1.00		C
ATOM	3290	CA			459	102.365	81.363	35.424	1.00		C
ATOM	3291	C			459	101.543	80.371	36.251 36.446	1.00		0
ATOM	3292	O CD			459	100.340	80.560	34.056	1.00		C
ATOM	3293	CB			459	102.725	80.779 80.716	32.938	1.00		C
ATOM	3294	CG			459	101.699			1.00		C
ATOM	3295 3296	CD1	LEU LEU		459 459	101.087 102.404	82.092 80.226	32.708 31.689	1.00		C
ATOM						102.404	79.322	36.754		30.22	N
ATOM	3297	N CA			460 460	102.190	78.322	37.561		30.19	C
ATOM	3298	CA C			460	101.483	78.973			30.56	C
ATOM	3299		HID	A	<del>1</del> 00	100.900	, 0 . 9 / 3	20.020	1.00		_



MOTA	3300	0	HIS	A	460	99.824	78.792	39.228	1.00	30.63	0
ATOM	3301	CB	HIS	Α	460	102.378	77.138	37.927	1.00	30.63	C
ATOM	3302	CG	HIS	Α	460	102.860	76.337	36.753	1.00	31.06	C
MOTA	3303	ND1	HIS	Α	460	102.178	76.270	35.558	1.00	31.79	N
ATOM	3304	CD2	HIS	Α	460	103.925	75.507	36.620	1.00	31.34	C
ATOM	3305	CE1	HIS	Α	460	102.802	75.437	34.742	1.00	31.58	C
ATOM	3306	NE2	HIS	A	460	103.865	74.959	35.361	1.00	30.54	N
MOTA	3307	N	GLU	Α	461	101.853	79.733	39.509	1.00	31.26	N
ATOM	3308	CA	GLU	Α	461	101.538	80.387	40.789	1.00	32.15	C
MOTA	3309	C	GLU	Α	461	100.548	81.562	40.795	1.00	30.88	C
MOTA	3310	0	GLU	Α	461	99.740	81.687	41.715	1.00	29.77	0
MOTA	3311	CB	GLU	A	461	102.833	80.887	41.457	1.00	33.27	C
MOTA	3312	CG	GLU	A	461	103.899	79.832	41.728	1.00	35.31	C
MOTA	3313	CD	GLU	Α	461	103.422	78.711	42.633	1.00	36.84	С
MOTA	3314	OE1	GLU	Α	461	102.952	78.986	43.758	1.00	37.87	0
ATOM	3315	OE2	GLU	Α	461	103.533	77.543	42.214	1.00	40.15	0
MOTA	3316	N	ASN	A	462	100.616	82.417	39.779	1.00	29.95	N
MOTA	3317	CA	ASN	Α	462	99.779	83.611	39.733	1.00	30.16	C
ATOM	3318	C	ASN	A	462	98.589	83.674	38.774	1.00	29.60	C
MOTA	3319	0	ASN	A	462	97.568	84.282	39.107	1.00	30.16	0
MOTA	3320	CB	ASN	A	462	100.683	84.817	39.484	1.00	31.63	C
ATOM	3321	CG	ASN	A	462	101.759	84.960	40.543	1.00	32.75	C
MOTA	3322	OD1	ASN	Α	462	102.725	85.696	40.367	1.00	35.69	0
MOTA	3323	ND2	ASN	Α	462	101.588	84.262	41.656	1.00	31.98	N
MOTA	3324	N	ALA	Α	463	98.711	83.066	37.598	1.00	27.52	N
MOTA	3325	CA	ALA	Α	463	97.634	83.103	36.610	1.00	26.51	C
MOTA	3326	C	ALA	A	463	96.338	82.451	37.076	1.00	25.73	C
MOTA	3327	0	ALA	Α	463	96.348	81.386	37.679	1.00	25.89	0
MOTA	3328	CB	ALA	Α	463	98.096	82.465	35.313	1.00	24.91	C
MOTA	3329	N	GLN	A	464	95.222	83.113	36.785		25.80	N
MOTA	3330	CA	GLN	A	464	93.888	82.619	37.136		25.60	C
MOTA	3331	С	GLN	A	464	92.959	82.601	35.923		24.30	C
MOTA	3332	0			464	93.147	83.369	34.977		23.88	0
MOTA	3333	CB			464	93.270	83.492	38.214		25.18	C
MOTA	3334	CG			464	93.996	83.413	39.517		28.22	C
ATOM	3335	CD			464	93.378	84.314	40.545	1.00		C
MOTA	3336	OE1			464	93.378	85.539	40.395	1.00		0
MOTA	3337	NE2			464	92.829	83.716	41.596		30.03	N
MOTA	3338	N			465	91.962	81.721	35.959	1.00		N
ATOM	3339	CA	PHE		465	91.009	81.614	34.868	1.00		C
ATOM	3340	С	PHE		465	89.627	82.121	35.246	1.00		C
ATOM	3341	0	PHE		465	89.199	82.035	36.404		23.64	0
ATOM	3342	CB			465	90.820	80.162	34.415		22.18	C
MOTA	3343	CG			465	92.042	79.513	33.836		23.68	C
ATOM	3344		PHE			92.775	78.594	34.577		25.25	C
MOTA	3345		PHE			92.437	79.780	32.533		23.42	C
ATOM	3346	CE1	PHE	A	465	93.884	77.947	34.025	1.00	24.49	C

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MOTA	3347	CE2	PHE	Α	465	93.544	79.138	31.974	1.00	23.33	C
ATOM	3348	CZ	PHE	A	465	94.265	78.220	32.724	1.00	24.38	C
ATOM	3349	N	VAL	A	466	88.932	82.656	34.251	1.00	22.63	N
ATOM	3350	CA	VAL	A	466	87.556	83.094	34.415	1.00	21.20	С
ATOM	3351	С	VAL	Α	466	86.828	82.188	33.423	1.00	21.50	C
ATOM	3352	0	VAL	A	466	87.281	81.994	32.300	1.00	19.95	0
ATOM	3353	CB	VAL	A	466	87.352	84.586	34.061	1.00	19.34	С
MOTA	3354	CG1	VAL	A	466	87.897	84.887	32.698	1.00	18.84	С
MOTA	3355	CG2	VAL	A	466	85.872	84.924	34.123	1.00	18.58	С
ATOM	3356	N	GLU	A	467	85.726	81.601	33.854	1.00	22.06	N
ATOM	3357	CA	GLU	A	467	84.979	80.698	33.005	1.00	24.06	C
ATOM	3358	C	GLU	A	467	84.050	81.439	32.068	1.00	24.75	С
MOTA	3359	0	GLU	A	467	83.415	82.413	32.465	1.00	25.25	0
ATOM	3360	CB	GLU	Α	467	84.151	79.763	33.858	1.00	24.21	C
MOTA	3361	CG	GLU	Α	467	83.546	78.653	33.077	1.00	28.44	С
ATOM	3362	CD	GLU	A	467	82.524	77.934	33.879	1.00	32.19	C
ATOM	3363	OE1	GLU	Α	467	82.756	77.771	35.093	1.00	34.37	0
MOTA	3364	OE2	GLU	A	467	81.502	77.521	33.298	1.00		0
HETATM	3365	N	MSE	A	468	83.956	80.981	30.826	1.00	24.94	N
HETATM	3366	CA	MSE	A	468	83.066	81.631	29.878	1.00		C
HETATM	3367	C	MSE	A	468	82.086	80.645	29.274	1.00	26.74	C
HETATM	3368	0	MSE	A	468	82.353	79.444	29.218	1.00		0
HETATM		CB			468	83.858	82.322	28.763		28.88	C
HETATM		CG			468	84.730	81.421	27.918		31.63	C
HETATM		SE			468	85.425	82.387	26.542		39.89	SE
HETATM		CE			468	86.595	81.206	25.795	1.00		C
ATOM	3373	N			469	80.945	81.156	28.827		27.65	N C
ATOM	3374	CA			469	79.919	80.314	28.212		28.52	C
ATOM	3375	C			469	80.148	80.275	26.705	1.00	29.26 28.60	0
ATOM	3376	O			469	81.093	80.886 80.888	26.197 28.488		27.30	C
ATOM	3377	CB			469	78.532	82.148	27.847		27.56	0
ATOM	3378	OG M			469	78.385 79.271	79.566	25.996		31.42	N
ATOM	3379	N			470 470	79.384	79.479	24.549		33.33	C
ATOM ATOM	3380 3381	CA C			470	79.362	80.860	23.920		35.22	Ċ
ATOM	3382	0			470	80.083	81.131	22.956		36.54	0
ATOM	3383	N			471	78.530	81.738	24.477		35.45	N
ATOM	3384	CA			471	78.408	83.106	23.992		36.13	C
ATOM	3385	C			471	79.720	83.867	24.206		36.97	C
ATOM	3386	0			471	80.182	84.595	23.321		37.26	. 0
ATOM	3387	СВ			471	77.268	83.809	24.722		35.98	С
ATOM	3388	N			472	80.311	83.701	25.388		37.72	N
ATOM	3389	CA			472	81.565	84.369	25.691		38.01	C
ATOM	3390	C			472	82.656	83.924	24.730		38.54	C
ATOM	3391	0			472	83.558	84.695	24.385		38.01	0
ATOM	3392	N			473	82.573	82.667	24.296		38.34	N
ATOM	3393	CA			473	83.550	82.125	23.363	1.00	37.79	С

#### LEU A 473 ATOM 3394 83.349 82.890 22.050 1.00 37.07 C MOTA 3395 **LEU A 473** 84.316 83.285 21.395 0 1.00 36.58 0 83.306 80.624 MOTA 3396 CB LEU A 473 23.152 C 1.00 38.74 3397 LEU A 473 MOTA CG 22.761 C 84.497 79.733 1.00 39.23 CD1 LEU A 473 83.986 78.347 MOTA 3398 22.459 1.00 38.60 C **ATOM** CD2 LEU A 473 80.274 21.558 C 3399 85.229 1.00 40.46 3400 ATOM N ILE A 474 82.084 83.108 21.687 N 1.00 36.82 **ATOM** 83.841 20.463 3401 CA ILE A 474 81.737 C 1.00 36.70 **ATOM** ILE A 474 82.348 85.238 3402 C 20.536 C 1.00 36.56 **ATOM** ILE A 474 0 19.582 3403 82.968 85.707 1.00 36.03 0 ILE A 474 80.198 84.001 20.294 MOTA 3404 CB 1.00 36.35 C 20.276 CG1 ILE A 474 1.00 36.33 C MOTA 3405 79.516 82.630 19.006 **ATOM** CG2 ILE A 474 79.889 84.760 C 3406 1.00 35.09 C **ATOM** 3407 CD1 ILE A 474 79.989 81.719 1.00 35.80 19.171 GLU A 475 85.899 21.674 1.00 35.95 ATOM 3408 N 82.159 N 3409 **GLU A 475** 82.700 87.233 MOTA CA C 21.880 1.00 36.17 **ATOM** C 84.216 87.235 3410 GLU A 475 21.791 1.00 36.53 C **ATOM** 3411 **GLU A 475** 84.807 88.209 21.327 1.00 36.42 0 0 23.254 ATOM 3412 CB **GLU A 475** 82.263 87.776 1.00 37.61 C 3413 **GLU A 475** 83.143 88.914 23.827 C MOTA CG 1.00 37.08 **GLU A 475** 82.626 C CD ATOM 3414 89.458 25.164 1.00 37.58 1.00 38.57 25.161 ATOM 3415 OE1 GLU A 475 81.614 90.188 0 3416 OE2 GLU A 475 83.216 89.146 26.221 0 ATOM 1.00 33.85 N SER A 476 ATOM 3417 84.841 86.145 22.237 1.00 37.04 N **ATOM** 3418 SER A 476 86.300 CA 86.039 22.242 C 1.00 37.27 85.973 20.851 1.00 37.06 C ATOM 3419 C SER A 476 86.924 **ATOM** SER A 476 88.046 86.442 20.638 3420 0 1.00 37.30 3421 CBSER A 476 86.728 84.830 C ATOM 23.068 1.00 36.80 84.963 SER A 476 24.391 1.00 36.30 ATOM 3422 OG 86.251 MOTA 3423 N HIS A 477 86.197 85.391 19.908 1.00 36.25 HIS A 477 3424 CA 86.667 85.288 18.533 ATOM 1.00 36.22 MOTA 3425 HIS A 477 86.181 86.477 C C 17.724 1.00 36.09 HIS A 477 MOTA 85.207 0 3426 87.141 18.100 1.00 34.72 C MOTA 3427 HIS A 477 86.132 84.008 17.897 CB1.00 36.36 C 3428 CG HIS A 477 86.901 MOTA 82.789 18.267 1.00 36.15 N ATOM 3429 ND1 HIS A 477 88.077 82.440 17.644 1.00 37.86 C CD2 HIS A 477 ATOM 86.690 3430 81.858 19.224 1.00 37.82 C ATOM CE1 HIS A 477 88.559 18.200 3431 81.343 1.00 38.68 NE2 HIS A 477 19.162 N ATOM 3432 87.736 80.969 1.00 38.42 PRO A 478 N 86.865 86.778 ATOM 3433 N 16.606 1.00 36.62 PRO A 478 C 86.409 MOTA 3434 CA 87.910 15.801 1.00 37.52 C MOTA 3435 C PRO A 478 84.974 87.589 15.387 1.00 38.47 84.626 0 86.417 15.192 MOTA 3436 0 PRO A 478 1.00 37.19 C ATOM 3437 PRO A 478 87.391 87.901 14.624 1.00 36.75 CBC MOTA 3438 CG PRO A 478 88.651 87.370 15.266 1.00 36.00 PRO A 478 15.987 C MOTA 3439 CD 88.056 86.173 1.00 36.59 HIS A 479 84.141 15.271 MOTA 3440 N 88.617 1.00 40.10



ATOM 3441 CA HIS A 479 82.749 88.409 14.898 1.00 41.71 ATOM 3442 C HIS A 479 82.661 89.607 14.178 1.00 42.96 ATOM 3443 CB HIS A 479 82.661 90.723 14.263 1.00 42.96 ATOM 3444 CB HIS A 479 81.906 88.138 16.142 1.00 42.96 ATOM 3445 CG HIS A 479 81.906 88.138 16.142 1.00 42.96 ATOM 3446 CD1 HIS A 479 83.105 89.373 17.989 1.00 44.02 ATOM 3447 CD2 HIS A 479 81.154 90.221 17.499 1.00 44.02 ATOM 3448 CB1 HIS A 479 82.292 90.423 18.773 1.00 43.65 ATOM 3449 NE2 HIS A 479 81.753 90.956 18.495 1.00 44.04 ATOM 3450 N ASP A 480 81.055 89.355 13.482 1.00 44.05 ATOM 3451 CA ASP A 480 81.355 89.355 13.482 1.00 44.05 ATOM 3452 C ASP A 480 81.196 91.161 11.787 1.00 45.86 ATOM 3453 C ASP A 480 81.196 91.161 11.787 1.00 45.86 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3456 CD1 ASP A 480 77.750 89.935 14.191 1.00 50.68 ATOM 3457 CD2 ASP A 480 77.750 89.935 14.191 1.00 50.68 ATOM 3458 N VAL A 481 82.987 91.093 10.141 1.00 50.68 ATOM 3458 N VAL A 481 82.987 91.093 10.141 1.00 45.97 ATOM 3460 C VAL A 481 82.983 90.657 11.099 1.00 46.00 ATOM 3461 C VAL A 481 83.437 90.069 9.105 1.00 45.34 ATOM 3462 CB VAL A 481 83.437 90.069 9.105 1.00 45.34 ATOM 3463 CG1 VAL A 481 83.437 90.069 9.105 1.00 45.43 ATOM 3465 CD1 A 482 83.481 89.073 11.769 1.00 45.54 ATOM 3466 CA GGI VAL A 481 83.437 90.069 9.105 1.00 45.80 ATOM 3467 CG GLN A 482 83.481 89.073 11.769 1.00 45.54 ATOM 3468 CG1 VAL A 481 83.437 90.069 9.105 1.00 45.80 ATOM 3467 CG GLN A 482 83.481 89.073 11.769 1.00 45.66 ATOM 3467 CG GLN A 482 83.481 89.073 11.769 1.00 45.66 ATOM 3467 C GLN A 482 83.491 99.035 5.466 1.00 45.66 ATOM 3467 C GLN A 482 83.491 99.035 5.466 1.00 45.66 ATOM 3467 C GLN A 482 83.496 90.00 6.353 1.00 41.65 ATOM 3468 C GLN A 482 83.492 89.494 93.190 1.00 45.80 ATOM 3470 CG GLN A 482 83.898 90.00 6.600 1.00 45.60 ATOM 3470 CG GLN A 482 83.898 90.00 6.600 1.00 45.60 ATOM 3470 CG GLN A 482 83.898 90.00 6.600 1.00 45.60 ATOM 3470 CG GLN A 482 83.129 90.035 5.666 1.00 45.60 ATOM 3470 CG GLN A 482 83.129 90.035 5.666 1.00 45.60 ATOM 3470 CG GLN A 482 83.129 9												
ATOM 3443 O HIS A 479 82.691 90.723 14.263 1.00 42.96 ATOM 3445 CB HIS A 479 81.906 88.138 16.142 1.00 42.96 ATOM 3445 CB HIS A 479 82.004 89.217 17.175 1.00 42.96 ATOM 3446 CB HIS A 479 83.105 89.373 17.989 1.00 44.02 ATOM 3447 CD2 HIS A 479 81.154 90.221 17.499 1.00 44.12 ATOM 3448 CBI HIS A 479 82.929 90.423 18.773 1.00 44.04 ATOM 3450 N ASP A 480 81.055 89.355 13.482 1.00 44.04 ATOM 3451 CA ASP A 480 80.321 90.373 12.741 1.00 45.86 ATOM 3451 CA ASP A 480 80.321 90.373 12.741 1.00 45.95 ATOM 3455 CG ASP A 480 81.055 89.355 13.482 1.00 45.95 ATOM 3455 CG ASP A 480 81.074 92.381 11.676 1.00 45.95 ATOM 3456 CD1 ASP A 480 79.631 91.326 13.716 1.00 45.95 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 779.631 91.326 13.716 1.00 47.55 ATOM 3456 CD1 ASP A 480 779.631 91.326 13.716 1.00 47.55 ATOM 3456 CD1 ASP A 480 779.631 91.326 13.716 1.00 47.55 ATOM 3456 CD1 ASP A 480 779.631 91.326 13.716 1.00 50.67 ATOM 3457 CD2 ASP A 480 779.631 91.326 13.716 1.00 50.67 ATOM 3458 N VAL A 481 82.083 90.601 14.672 1.00 50.67 ATOM 3458 C VAL A 481 82.083 90.457 11.099 1.00 45.95 ATOM 3459 CA VAL A 481 82.083 90.457 11.099 1.00 45.95 ATOM 3461 C VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3462 CB VAL A 481 83.781 88.934 9.438 1.00 45.58 ATOM 3462 CB VAL A 481 83.781 88.934 9.438 1.00 45.54 ATOM 3465 N GIN A 482 83.841 90.475 7.482 1.00 45.62 ATOM 3465 N GIN A 482 83.842 99.600 6.756 1.00 45.56 ATOM 3467 C GIN A 482 83.842 99.600 6.756 1.00 45.56 ATOM 3467 C GIN A 482 83.842 99.600 6.756 1.00 45.50 ATOM 3467 C GIN A 482 83.842 99.600 6.756 1.00 45.50 ATOM 3470 C GIN A 482 83.822 89.600 6.756 1.00 45.62 ATOM 3470 C GIN A 482 83.822 89.600 6.756 1.00 45.62 ATOM 3470 C GIN A 482 83.826 89.807 6.600 1.00 45.50 ATOM 3470 C GIN A 482 83.846 89.477 1.760 1.00 45.40 ATOM 3470 C GIN A 482 83.846 89.477 1.760 1.00 45.40 ATOM 3470 C GIN A 482 83.846 89.479 1.900 4.00 45.60 ATOM 3470 C GIN A 482 83.86 89.479 1.900 1.00 45.60 ATOM 3470 C GIN A 482 83.86 89.877 87.626 5.378 1.00 46.60 ATOM 3470 C GIN A 482 83.86	ATOM :	3441	CA	HIS	A	479	82.749	88.409	14.898	1.00	41.71	C
ATOM         3444         CB         HIS A 479         81.906         88.138         16.142         1.00         42.19           ATOM         3445         NDI         HIS A 479         82.004         89.217         17.175         1.00         42.96           ATOM         3446         NDI         HIS A 479         81.154         90.221         17.499         1.00         44.12           ATOM         3448         CE1         HIS A 479         81.154         90.221         17.499         1.00         44.12           ATOM         3448         CE1         HIS A 479         81.753         90.955         18.495         1.00         44.04           ATOM         3450         N         ASP A 480         81.055         89.355         13.482         1.00         44.35           ATOM         3451         CA         ASP A 480         81.074         92.381         11.676         1.00         45.86           ATOM         3454         CB         ASP A 480         79.631         91.326         13.716         1.00         50.07           ATOM         3455         OD2         ASP A 480         77.750         89.935         14.191         1.00         50.68 <td>MOTA</td> <td>3442</td> <td>C</td> <td>HIS</td> <td>A</td> <td>479</td> <td>82.163</td> <td>89.607</td> <td>14.178</td> <td>1.00</td> <td>43.00</td> <td>C</td>	MOTA	3442	C	HIS	A	479	82.163	89.607	14.178	1.00	43.00	C
ATOM 3445 CG HIS A 479 82.004 89.217 17.175 1.00 42.96 ATOM 3446 ND1 HIS A 479 83.105 89.373 17.899 1.00 44.02 ATOM 3447 CD2 HIS A 479 83.105 89.373 17.899 1.00 44.02 ATOM 3449 NEZ HIS A 479 82.929 90.423 18.773 1.00 43.65 ATOM 3449 NEZ HIS A 479 81.753 90.956 18.495 1.00 44.04 ATOM 3450 N ASP A 480 81.055 89.355 13.482 1.00 44.35 ATOM 3451 CA ASP A 480 81.055 89.355 13.482 1.00 44.35 ATOM 3452 C ASP A 480 81.196 91.161 11.787 1.00 45.86 ATOM 3453 O ASP A 480 81.076 91.161 11.787 1.00 45.86 ATOM 3453 O ASP A 480 81.076 91.161 11.787 1.00 45.86 ATOM 3455 CG ASP A 480 81.076 91.161 11.787 1.00 45.86 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 46.64 ATOM 3456 ODI ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3456 ODI ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3456 ODI ASP A 480 78.698 90.601 14.672 1.00 50.07 ATOM 3456 ODI ASP A 480 78.912 90.698 15.904 1.00 52.11 ATOM 3458 N VAL A 481 82.083 90.457 11.099 1.00 45.97 ATOM 3450 CC VAL A 481 82.975 91.093 10.141 1.00 45.97 ATOM 3460 C VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3461 O VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3463 CGI VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3463 CGI VAL A 481 84.199 91.748 10.861 1.00 45.54 ATOM 3463 CGI VAL A 481 84.199 91.748 10.861 1.00 45.54 ATOM 3466 CG VAL A 481 84.199 91.748 10.861 1.00 45.54 ATOM 3466 CG CG VAL A 481 84.199 91.748 10.861 1.00 45.54 ATOM 3466 CG CG CLN A 482 83.812 89.600 6.756 1.00 45.66 ATOM 3467 C CG CLN A 482 83.812 89.600 6.756 1.00 45.66 ATOM 3467 C CG CLN A 482 83.812 89.600 6.756 1.00 45.66 ATOM 3467 C CG CLN A 482 83.812 89.600 6.756 1.00 45.66 ATOM 3470 CG CLN A 482 83.826 89.447 3.190 1.00 45.54 ATOM 3470 CG CLN A 482 83.826 89.447 3.190 1.00 45.57 ATOM 3472 CG CLN A 482 83.826 89.447 3.190 1.00 45.50 ATOM 3473 NC2 CG CLN A 482 83.826 89.447 3.190 1.00 45.50 ATOM 3473 NC2 CG CLN A 482 83.826 89.447 3.190 1.00 46.60 ATOM 3476 C C CLN A 483 87.431 88.406 6.643 1.00 46.60 ATOM 3476 C C CLN A 483 87.431 88.406 6.643 1.00 46.60 ATOM 3477 C C CLL A 483 87.437 88.500 6.760 1.00 47.02	MOTA	3443	0	HIS	Α	479	82.691	90.723	14.263	1.00	42.96	0
ATOM 3446 ND1 HIS A 479 83.105 89.373 17.989 1.00 44.02 ATOM 3447 CD2 HIS A 479 81.154 90.221 17.499 1.00 44.02 ATOM 3448 CE1 HIS A 479 81.753 90.423 18.773 1.00 43.65 ATOM 3449 NE2 HIS A 479 81.753 90.956 18.495 1.00 44.04 ATOM 3450 N ASP A 480 81.055 89.355 13.482 1.00 44.05 ATOM 3451 CA ASP A 480 81.055 89.355 13.482 1.00 45.86 ATOM 3451 CA ASP A 480 81.196 91.161 11.787 1.00 45.86 ATOM 3453 O ASP A 480 81.074 92.381 11.676 1.00 45.86 ATOM 3455 CG ASP A 480 81.074 92.381 11.676 1.00 46.64 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 77.750 89.935 14.191 1.00 50.07 ATOM 3456 ODI ASP A 480 77.750 89.935 14.191 1.00 50.68 ATOM 3457 ODZ ASP A 480 78.698 90.601 14.672 1.00 50.07 ATOM 3458 N VAL A 481 82.983 90.457 11.099 1.00 46.00 ATOM 3459 CA VAL A 481 82.983 90.457 11.099 1.00 45.86 ATOM 3450 C VAL A 481 83.781 88.934 9.1041 1.00 45.97 ATOM 3460 C VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3462 CB VAL A 481 83.781 88.934 9.438 1.00 46.24 ATOM 3465 CG VAL A 481 83.781 88.934 9.388 1.00 44.87 ATOM 3465 CG VAL A 481 85.177 92.301 9.834 1.00 46.24 ATOM 3465 CG VAL A 481 85.177 92.301 9.834 1.00 45.62 ATOM 3466 CA GLN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLN A 482 83.822 89.0706 6.553 1.00 45.62 ATOM 3466 CA GLN A 482 83.812 89.0706 6.553 1.00 45.62 ATOM 3467 C GLN A 482 83.812 89.894 9.733 1.00 44.56 ATOM 3467 C GLN A 482 83.812 89.894 9.706 6.553 1.00 45.02 ATOM 3471 CD GLN A 482 83.812 89.894 9.706 6.553 1.00 45.02 ATOM 3476 C GLN A 482 83.812 89.894 9.706 6.553 1.00 45.02 ATOM 3476 C GLN A 482 83.812 89.894 9.706 6.553 1.00 45.02 ATOM 3477 CD GLN A 482 83.822 89.0706 6.556 1.00 45.02 ATOM 3476 C GLN A 482 83.812 89.894 9.706 6.503 1.00 45.00 ATOM 3476 C GLN A 482 83.894 89.130 4.279 1.00 45.02 ATOM 3476 C GLN A 483 87.797 88.500 6.766 1.00 45.00 ATOM 3476 C GLN A 483 87.797 87.626 5.378 1.00 47.02 ATOM 3478 C GLL A 483 87.797 87.626 5.378 1.00 47.02 ATOM	MOTA	3444	CB	HIS	A	479	81.906	88.138	16.142	1.00	42.19	C
ATOM 3447 CD2 HIS A 479 81.154 90.221 17.499 1.00 44.12 ATOM 3448 CE1 HIS A 479 82.929 90.423 18.773 1.00 43.65 ATOM 3449 NE2 HIS A 479 81.755 90.956 18.495 1.00 44.04 ATOM 3450 N ASP A 480 81.055 89.355 13.482 1.00 44.35 ATOM 3451 CA ASP A 480 81.055 89.355 13.482 1.00 44.35 ATOM 3451 CA ASP A 480 81.965 91.161 11.787 1.00 45.95 ATOM 3454 CB ASP A 480 81.074 92.381 11.676 1.00 46.64 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 77.750 89.935 14.191 1.00 50.07 ATOM 3455 CG ASP A 480 77.750 89.935 14.191 1.00 50.08 ATOM 3457 OD2 ASP A 480 77.750 89.935 14.191 1.00 50.08 ATOM 3458 N VAL A 481 82.083 90.457 11.099 1.00 46.00 ATOM 3458 N VAL A 481 82.083 90.457 11.099 1.00 46.00 ATOM 3450 C VAL A 481 82.083 90.457 11.099 1.00 46.00 ATOM 3460 C VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3461 O VAL A 481 84.199 91.748 10.861 1.00 44.87 ATOM 3463 CG1 VAL A 481 84.199 91.748 10.861 1.00 45.58 ATOM 3463 CG1 VAL A 481 84.199 91.748 10.861 1.00 45.58 ATOM 3466 CG VAL A 481 84.199 91.748 10.861 1.00 45.58 ATOM 3466 CG2 VAL A 481 84.199 91.748 10.861 1.00 45.58 ATOM 3466 CG2 VAL A 481 84.199 91.748 10.861 1.00 45.58 ATOM 3466 CG CGL VAL A 481 84.199 91.748 10.861 1.00 45.58 ATOM 3466 CG CGL VAL A 481 84.199 91.748 10.861 1.00 45.56 ATOM 3466 CG CGL VAL A 482 83.416 90.475 7.842 1.00 45.66 ATOM 3466 CG CGL VAL A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3467 C CGLN A 482 83.819 90.706 6.353 1.00 44.56 ATOM 3467 C CGLN A 482 83.825 89.800 6.766 1.00 45.04 ATOM 3470 CG GLN A 482 83.826 89.600 6.756 1.00 45.04 ATOM 3470 CG GLN A 482 83.826 89.600 6.756 1.00 45.04 ATOM 3470 CG GLN A 482 83.826 89.600 6.756 1.00 45.04 ATOM 3470 CG GLN A 482 83.894 89.493 3.095 1.00 45.00 ATOM 3470 CG GLN A 482 83.894 89.493 3.095 1.00 45.00 ATOM 3470 CG GLN A 482 83.895 89.600 6.766 1.00 45.00 ATOM 3470 CG GLN A 482 83.896 80.497 91.00 40.00	MOTA	3445	CG	HIS	A	479	82.004	89.217	17.175	1.00	42.96	C
ATOM 3448 CE1 HIS A 479	MOTA	3446	ND1	HIS	A	479	83.105	89.373	17.989	1.00	44.02	N
ATOM 3449 NE2 HIS A 479	MOTA	3447	CD2	HIS	A	479	81.154	90.221	17.499	1.00	44.12	C
ATOM 3450 N ASP A 480 81.055 89.355 13.482 1.00 44.35 ATOM 3451 CA ASP A 480 80.321 90.373 12.741 1.00 45.86 ATOM 3453 C ASP A 480 81.196 91.161 11.787 1.00 45.86 ATOM 3453 O ASP A 480 81.074 92.381 11.676 1.00 46.64 ATOM 3454 CB ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 78.698 90.601 14.672 1.00 50.07 ATOM 3456 OD1 ASP A 480 77.750 89.935 14.191 1.00 50.68 ATOM 3456 OD1 ASP A 480 77.750 89.935 14.191 1.00 50.68 ATOM 3456 OD1 ASP A 480 77.750 89.935 14.191 1.00 52.11 ATOM 3458 N VAL A 481 82.975 91.093 10.141 1.00 45.97 ATOM 3458 N VAL A 481 82.975 91.093 10.141 1.00 45.97 ATOM 3450 C VAL A 481 82.975 91.093 10.141 1.00 45.97 ATOM 3460 C VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3461 O VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 45.33 ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.62 ATOM 3466 CA GLN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLN A 482 83.812 89.600 6.756 1.00 45.62 ATOM 3466 CA GLN A 482 83.812 89.600 6.756 1.00 45.62 ATOM 3466 CB GLN A 482 85.831 89.650 6.600 1.00 45.00 ATOM 3467 C GLN A 482 83.812 89.600 6.756 1.00 45.00 ATOM 3467 C GLN A 482 85.831 89.650 6.600 1.00 45.00 ATOM 3470 CG GLN A 482 83.312 90.035 5.466 1.00 45.00 ATOM 3471 CD GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 83.384 89.49 3.095 1.00 45.00 ATOM 3472 CE GLN A 482 83.384 89.49 3.095 1.00 45.00 ATOM 3473 NE2 GLN A 482 83.384 89.49 3.095 1.00 45.00 ATOM 3473 NE2 GLN A 482 83.384 89.49 3.095 1.00 45.00 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3473 NE2 GLN A 482 83.384 89.49 3.095 1.00 45.00 ATOM 3473 NE2 GLN A 482 83.384 89.49 3.095 1.00 47.00 45.00 ATOM 3473 NE2 GLN A 482 83.384 89.49 3.095 1.00 45.00 ATOM 3473 NE2 GLN A 482 83.384 89.49 3.095 1.00 47.00 45.00 ATOM 3474 N ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 45.70 ATOM 3474 C ILE A 483 87.797 87.626 5.378 1.00 47.10 47.02 ATOM 3478 CB ILE A 483 87.797 87.626 5.378 1.00 47.10 47.20 ATOM 3484 C ILE A 483	MOTA	3448	CE1	HIS	A	479	82.929	90.423	18.773	1.00	43.65	C
ATOM 3451 CA ASP A 480 80.321 90.373 12.741 1.00 45.86 ATOM 3452 C ASP A 480 81.196 91.161 11.787 1.00 45.95 ATOM 3453 O ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 47.55 ATOM 3455 CG ASP A 480 78.698 90.601 14.672 1.00 50.68 ATOM 3455 ODL ASP A 480 78.912 90.698 15.904 1.00 50.68 ATOM 3455 ODL ASP A 480 78.912 90.698 15.904 1.00 52.11 ATOM 3456 ODL ASP A 481 82.083 90.457 11.099 1.00 46.00 ATOM 3458 N VAL A 481 82.083 90.457 11.099 1.00 45.97 ATOM 3450 C VAL A 481 83.437 90.069 9.105 1.00 44.87 ATOM 3450 C VAL A 481 83.437 90.069 9.105 1.00 44.87 ATOM 3461 O VAL A 481 83.437 90.069 9.105 1.00 44.87 ATOM 3463 CGI VAL A 481 84.199 91.748 10.861 1.00 45.54 ATOM 3465 N GLA A 481 84.199 91.748 10.861 1.00 45.53 ATOM 3466 CG VAL A 481 84.199 91.748 10.861 1.00 45.63 ATOM 3465 N GLA A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CG CG VAL A 481 85.177 92.301 9.834 1.00 45.63 ATOM 3466 CG CG VAL A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CG GLA A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3466 CG GLA A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3468 O GLA A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3469 CG GLA A 482 83.829 89.600 6.756 1.00 44.56 ATOM 3467 C GLA A 482 83.829 89.600 6.756 1.00 45.66 ATOM 3467 C GLA A 482 83.829 89.600 6.756 1.00 45.66 ATOM 3467 C GLA A 482 83.829 89.600 6.756 1.00 45.66 ATOM 3470 CG GLA A 482 83.829 89.600 6.756 1.00 45.66 ATOM 3470 CG GLA A 482 83.829 89.600 6.756 1.00 45.66 ATOM 3473 NEZ GLA A 482 83.829 89.600 6.756 1.00 45.66 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 1.00 44.570 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 1.00 47.02 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 1.00 47.02 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 1.00 47.02 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 1.00 47.02 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 1.00 47.02 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 70 1.00 47.02 ATOM 3473 NEZ GLA A 483 85.977 88.500 6.766 70 1.00 47.02 ATOM 3473 NEZ GLA A 483 88.507 87.668 7.721 7.804 1.00 47.02 ATOM 3480 CG IL	MOTA	3449	NE2	HIS	A	479	81.753	90.956	18.495	1.00	44.04	N
ATOM 3452 C ASP A 480 81.196 91.161 11.787 1.00 45.95 ATOM 3453 O ASP A 480 81.074 92.381 11.676 1.00 46.64 ATOM 3455 CG ASP A 480 79.631 91.326 13.716 1.00 45.95 ATOM 3456 CG ASP A 480 78.698 90.601 14.672 1.00 50.07 ATOM 3456 OD1 ASP A 480 77.750 89.935 14.191 1.00 50.07 ATOM 3457 OD2 ASP A 480 78.698 90.601 14.672 1.00 50.07 ATOM 3458 N VAL A 481 82.083 90.457 11.099 1.00 46.00 ATOM 3459 CA VAL A 481 82.083 90.457 11.099 1.00 45.97 ATOM 3450 CA VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3461 O VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 46.24 ATOM 3463 CGI VAL A 481 84.199 91.748 10.861 1.00 46.24 ATOM 3465 N GLI A 481 85.177 92.301 9.834 1.00 45.43 ATOM 3465 CA GLIN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLIN A 482 83.416 90.475 7.842 1.00 45.66 ATOM 3466 CA GLIN A 482 83.416 90.475 7.842 1.00 45.66 ATOM 3467 CG GLIN A 482 83.8129 90.035 5.466 1.00 45.64 ATOM 3467 CG GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3467 CG GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3470 CG GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3470 CG GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3470 CG GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3473 NE2 GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3473 NE2 GLIN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3473 NE2 GLIN A 482 83.126 89.487 1.976 1.00 53.04 ATOM 3473 NE2 GLIN A 482 83.126 89.487 1.976 1.00 53.04 ATOM 3473 NE2 GLIN A 482 83.126 89.487 1.976 1.00 53.04 ATOM 3473 NE2 GLIN A 483 85.777 88.500 6.760 1.00 45.40 ATOM 3474 N ILE A 483 87.797 88.500 6.760 1.00 45.80 ATOM 3475 CG ILE A 483 87.797 88.500 6.760 1.00 45.40 ATOM 3475 CG ILE A 483 87.797 88.500 6.760 1.00 47.02 ATOM 3478 CG ILE A 483 87.797 88.500 6.760 1.00 47.02 ATOM 3478 CG ILE A 483 88.710 87.797 87.626 5.378 1.00 46.80 ATOM 3479 CG ILE A 483 88.710 87.797 87.626 5.378 1.00 47.02 ATOM 3478 CG ILE A 483 88.573 88.240 4.491 1.00 47.02 ATOM 3480 CG ILE A 483 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM	MOTA	3450	N	ASP	A	480	81.055	89.355	13.482	1.00	44.35	N
ATOM 3453 O ASP A 480	MOTA	3451	CA	ASP	A	480	80.321	90.373	12.741	1.00	45.86	C
ATOM 3454 CB ASP A 480	MOTA	3452	C	ASP	A	480	81.196	91.161	11.787	1.00	45.95	C
ATOM 3455 CG ASP A 480	MOTA	3453	0	ASP	Α	480	81.074	92.381	11.676	1.00	46.64	0
ATOM 3456 OD1 ASP A 480	MOTA	3454	CB	ASP	Α	480	79.631	91.326	13.716	1.00	47.55	C
ATOM 3457 OD2 ASP A 480	MOTA	3455	CG	ASP	A	480	78.698	90.601	14.672	1.00	50.07	C
ATOM 3458 N VAL A 481 82.083 90.457 11.099 1.00 46.00 ATOM 3459 CA VAL A 481 82.975 91.093 10.141 1.00 45.97 ATOM 3460 C VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3461 O VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 46.24 ATOM 3463 CGI VAL A 481 84.888 90.733 11.769 1.00 46.53 ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.43 ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.43 ATOM 3466 CA GLN A 482 83.416 90.475 7.842 1.00 45.66 ATOM 3466 CA GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 45.54 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 83.820 89.447 3.190 1.00 51.02 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3476 C ILE A 483 85.777 88.500 6.760 1.00 45.70 ATOM 3476 C ILE A 483 87.479 88.406 6.643 1.00 46.80 ATOM 3477 O ILE A 483 87.479 88.406 6.643 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3480 CG2 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3481 CD1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3481 CD1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3482 CG ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.26 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.66 ATOM 3485 C THR A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 C THR A 484 88.573 88.250 6.599 1.00 47.69	MOTA	3456	OD1	ASP	A	480	77.750	89.935	14.191	1.00	50.68	0
ATOM 3459 CA VAL A 481 82.975 91.093 10.141 1.00 45.97  ATOM 3460 C VAL A 481 83.437 90.069 9.105 1.00 45.58  ATOM 3461 O VAL A 481 83.781 88.934 9.438 1.00 44.87  ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 46.24  ATOM 3463 CG1 VAL A 481 85.177 92.301 9.834 1.00 45.53  ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.43  ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.62  ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.62  ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.63  ATOM 3468 O GLN A 482 85.331 89.0706 6.353 1.00 44.56  ATOM 3469 CB GLN A 482 83.384 89.130 4.279 1.00 49.29  ATOM 3471 CD GLN A 482 83.384 89.130 4.279 1.00 49.29  ATOM 3472 OE1 GLN A 482 83.126 89.489 3.095 1.00 51.02  ATOM 3473 NE2 GLN A 482 83.126 89.487 1.976 1.00 50.81  ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70  ATOM 3475 CA ILE A 483 85.977 88.500 6.760 1.00 45.70  ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80  ATOM 3477 O ILE A 483 88.979 87.626 5.378 1.00 46.80  ATOM 3479 CG1 ILE A 483 87.797 87.626 5.378 1.00 46.80  ATOM 3479 CG1 ILE A 483 88.7.797 87.626 5.378 1.00 46.80  ATOM 3479 CG1 ILE A 483 88.7.797 87.626 5.378 1.00 46.80  ATOM 3479 CG1 ILE A 483 88.7.797 87.626 5.378 1.00 46.80  ATOM 3479 CG1 ILE A 483 88.7.565 88.314 9.172 1.00 47.22  ATOM 3480 CG2 ILE A 483 88.045 87.668 7.870 1.00 46.80  ATOM 3480 CG2 ILE A 483 88.045 87.668 7.870 1.00 47.22  ATOM 3480 CG2 ILE A 483 88.045 87.661 10.430 1.00 47.22  ATOM 3481 CD1 ILE A 483 88.045 87.661 10.430 1.00 47.26  ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.02  ATOM 3485 O THR A 484 88.570 87.655 3.258 1.00 47.69  ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 47.04	MOTA	3457	OD2	ASP	A	480	78.912	90.698	15.904	1.00	52.11	0
ATOM 3460 C VAL A 481 83.437 90.069 9.105 1.00 45.58 ATOM 3461 O VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 46.24 ATOM 3463 CG1 VAL A 481 84.888 90.733 11.769 1.00 46.53 ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.43 ATOM 3466 CA GLN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.62 ATOM 3466 CA GLN A 482 85.898 90.706 6.353 1.00 45.60 ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 82.506 89.487 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3475 CA ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3476 C ILE A 483 87.431 88.406 6.643 1.00 46.80 ATOM 3477 O ILE A 483 87.431 88.406 6.643 1.00 46.80 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.22 ATOM 3479 CG1 ILE A 483 87.378 86.479 5.209 1.00 47.22 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3478 CB ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3481 CD1 ILE A 483 88.573 88.240 4.491 1.00 47.22 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 CB ILE A 483 88.573 88.240 4.491 1.00 47.02 ATOM 3485 CB ILE A 483 88.573 88.240 4.491 1.00 47.02 ATOM 3485 CB ILE A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 CB ILE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.491 1.00 47.46 ATOM 3485 CB ITE A 484 88.573 88.240 4.291 1	MOTA	3458	N	VAL	A	481	82.083	90.457	11.099	1.00	46.00	N
ATOM 3461 O VAL A 481 83.781 88.934 9.438 1.00 44.87 ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 46.24 ATOM 3463 CG1 VAL A 481 85.177 92.301 9.834 1.00 45.53 ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.66 ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 45.64 ATOM 3470 CG GLN A 482 83.824 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OEI GLN A 482 83.126 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3475 CA ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3476 C ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3477 O ILE A 483 87.797 87.626 5.378 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.797 87.626 5.378 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.797 87.626 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 47.22 ATOM 3480 CG2 ILE A 483 88.557 88.240 4.491 1.00 47.22 ATOM 3481 CD1 ILE A 483 88.573 88.240 4.491 1.00 47.02 ATOM 3482 N THR A 484 88.870 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.8710 88.453 2.002 1.00 47.46 ATOM 3485 O THR A 484 88.8710 88.453 2.002 1.00 47.46	MOTA	3459	CA	VAL	Α	481	82.975	91.093	10.141	1.00	45.97	C
ATOM 3462 CB VAL A 481 84.199 91.748 10.861 1.00 46.24 ATOM 3463 CG1 VAL A 481 84.888 90.733 11.769 1.00 46.53 ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.43 ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.60 ATOM 3469 CB GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3472 OE1 GLN A 482 83.128 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 53.04 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3476 C ILE A 483 87.431 88.406 6.643 1.00 46.80 ATOM 3477 O ILE A 483 87.431 88.406 6.643 1.00 46.80 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.86 ATOM 3481 CD1 ILE A 483 88.045 87.668 7.870 1.00 47.22 ATOM 3481 CD1 ILE A 483 88.045 87.668 7.870 1.00 47.22 ATOM 3481 CD1 ILE A 483 88.573 88.240 4.491 1.00 47.26 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.26 ATOM 3485 O THR A 484 88.570 87.661 1.00 47.46 ATOM 3485 O THR A 484 88.570 87.665 2.559 1.00 47.46 ATOM 3485 O THR A 484 88.571 88.453 2.002 1.00 47.46 ATOM 3485 O THR A 484 88.571 88.453 2.002 1.00 47.46 ATOM 3485 O THR A 484 88.571 88.453 2.002 1.00 47.46	MOTA	3460	C	VAL	A	481	83.437	90.069	9.105	1.00	45.58	C
ATOM 3463 CG1 VAL A 481 84.888 90.733 11.769 1.00 46.53 ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.43 ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3468 O GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3469 CB GLN A 482 83.822 90.706 6.353 1.00 44.56 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 53.04 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3477 O ILE A 483 87.431 88.406 6.643 1.00 46.80 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.22 ATOM 3478 CB ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3481 CDI ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3481 CDI ILE A 483 88.045 87.668 7.870 1.00 47.22 ATOM 3481 CDI ILE A 483 88.045 87.664 1.00 46.86 ATOM 3483 CDI ILE A 483 88.045 87.664 1.00 47.22 ATOM 3481 CDI ILE A 483 88.045 87.664 1.00 47.22 ATOM 3483 CDI ILE A 483 88.045 87.664 1.00 47.22 ATOM 3483 CDI ILE A 483 88.045 87.664 1.00 47.22 ATOM 3483 CDI ILE A 483 88.573 88.240 4.91 1.00 47.26 ATOM 3484 CDI ILE A 483 88.573 88.240 4.91 1.00 47.26 ATOM 3483 CDI ILE A 484 88.573 88.240 4.91 1.00 47.26 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.26 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.26 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 4.91 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 5.565 3.258 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 5.565 3.258 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 5.565 3.258 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 5.565 3.258 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.240 5.565 3.258 1.00 47.66 ATOM 3485 CDI ILE A 484 88.573 88.24	MOTA	3461	0	VAL	A	481	83.781	88.934	9.438	1.00	44.87	0
ATOM 3464 CG2 VAL A 481 85.177 92.301 9.834 1.00 45.43 ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.62 ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3468 O GLN A 482 85.888 90.706 6.353 1.00 44.56 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 83.1280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 887.378 86.479 5.209 1.00 47.10 ATOM 3479 CG1 ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3480 CG2 ILE A 483 88.045 87.668 7.870 1.00 47.22 ATOM 3481 CD1 ILE A 483 88.956 87.721 7.804 1.00 47.22 ATOM 3482 N THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.46 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.46 ATOM 3485 O THR A 484 88.970 87.565 3.259 1.00 47.46 ATOM 3485 O THR A 484 88.970 87.565 3.259 1.00 47.46 ATOM 3485 O THR A 484 88.970 87.565 3.259 1.00 47.46	MOTA	3462	CB	VAL	Α	481	84.199	91.748	10.861	1.00	46.24	C
ATOM 3465 N GLN A 482 83.416 90.475 7.842 1.00 45.62  ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.66  ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.00  ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56  ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 46.54  ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29  ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02  ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04  ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81  ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70  ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40  ATOM 3476 C ILE A 483 87.378 86.479 5.209 1.00 47.10  ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10  ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 46.86  ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15  ATOM 3481 CD1 ILE A 483 88.045 87.668 7.870 1.00 46.15  ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02  ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.02  ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69  ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.46  ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 47.46	MOTA	3463	CG1	VAL	A	481	84.888	90.733	11.769	1.00	46.53	С
ATOM 3466 CA GLN A 482 83.822 89.600 6.756 1.00 45.66 ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 46.54 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 83.126 89.847 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 88.045 87.668 7.870 1.00 47.22 ATOM 3480 CG2 ILE A 483 88.566 87.721 7.804 1.00 47.22 ATOM 3481 CD1 ILE A 483 88.573 88.240 4.491 1.00 47.26 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3484 C THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3464	CG2	VAL	Α	481	85.177	92.301	9.834	1.00	45.43	C
ATOM 3467 C GLN A 482 85.331 89.650 6.600 1.00 45.00 ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 46.54 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.045 87.661 10.430 1.00 47.22 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.26 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.26 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3486 CB THR A 484 88.970 88.453 2.002 1.00 48.12	MOTA	3465	N	GLN	Α	482	83.416	90.475	7.842	1.00	45.62	N
ATOM 3468 O GLN A 482 85.898 90.706 6.353 1.00 44.56 ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 46.54 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.075 87.641 10.430 1.00 47.26 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.26 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 88.710 88.453 2.002 1.00 47.46 ATOM 3485 O THR A 484 88.710 88.453 2.002 1.00 47.46	MOTA	3466	CA	GLN	A	482	83.822	89.600	6.756	1.00	45.66	С
ATOM 3469 CB GLN A 482 83.129 90.035 5.466 1.00 46.54 ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 47.46 ATOM 3485 O THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3467	C	GLN	A	482	85.331	89.650	6.600	1.00	45.00	C
ATOM 3470 CG GLN A 482 83.384 89.130 4.279 1.00 49.29 ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 48.12	MOTA	3468	0	GLN	Α	482	85.898	90.706	6.353	1.00	44.56	0
ATOM 3471 CD GLN A 482 82.506 89.489 3.095 1.00 51.02 ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.573 88.240 4.491 1.00 47.69 ATOM 3484 C THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 47.46 ATOM 3485 O THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3469	CB	GLN	A	482	83.129	90.035	5.466	1.00	46.54	C
ATOM 3472 OE1 GLN A 482 81.280 89.447 3.190 1.00 53.04 ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3470	CG	GLN	Α	482	83.384	89.130	4.279	1.00	49.29	C
ATOM 3473 NE2 GLN A 482 83.126 89.847 1.976 1.00 50.81 ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3471	CD	GLN	Α	482	82.506	89.489	3.095	1.00	51.02	C
ATOM 3474 N ILE A 483 85.977 88.500 6.760 1.00 45.70 ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3472	OE1	GLN	Α	482	81.280	89.447	3.190	1.00	53.04	0
ATOM 3475 CA ILE A 483 87.431 88.406 6.643 1.00 46.40 ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3473	NE2	GLN	Α	482	83.126	89.847	1.976	1.00	50.81	N
ATOM 3476 C ILE A 483 87.797 87.626 5.378 1.00 46.80 ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10 ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3474	N	ILE	A	483	85.977	88.500	6.760	1.00	45.70	N
ATOM 3477 O ILE A 483 87.378 86.479 5.209 1.00 47.10  ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86  ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22  ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15  ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26  ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02  ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69  ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46  ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97  ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3475	CA	ILE	A	483	87.431	88.406	6.643	1.00	46.40	C
ATOM 3478 CB ILE A 483 88.045 87.668 7.870 1.00 46.86 ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3476	C	ILE	Α	483	87.797	87.626	5.378	1.00	46.80	C
ATOM 3479 CG1 ILE A 483 87.565 88.314 9.172 1.00 47.22 ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3477	Ο.	ILE	Α	483	87.378	86.479	5.209	1.00	47.10	0
ATOM 3480 CG2 ILE A 483 89.566 87.721 7.804 1.00 46.15 ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3478	CB	ILE	Α	483	88.045	87.668	7.870	1.00	46.86	C
ATOM 3481 CD1 ILE A 483 88.107 87.641 10.430 1.00 47.26 ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	ATOM	3479	CG1	ILE	Α	483	87.565	88.314	9.172	1.00	47.22	C
ATOM 3482 N THR A 484 88.573 88.240 4.491 1.00 47.02 ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3480	CG2	ILE	Α	483	89.566	87.721	7.804	1.00	46.15	C
ATOM 3483 CA THR A 484 88.970 87.565 3.258 1.00 47.69 ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3481	CD1	ILE	Α	483	88.107	87.641	10.430	1.00	47.26	C
ATOM 3484 C THR A 484 90.448 87.192 3.316 1.00 47.46 ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3482	N	THR	A	484	88.573	88.240	4.491	1.00	47.02	N
ATOM 3485 O THR A 484 90.908 86.296 2.599 1.00 46.97 ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3483	CA	THR	A	484	88.970	87.565	3.258	1.00	47.69	C
ATOM 3486 CB THR A 484 88.710 88.453 2.002 1.00 48.12	MOTA	3484	C	THR	Α	484	90.448	87.192	3.316	1.00	47.46	C
	MOTA	3485	0	THR	Α	484	90.908	86.296	2.599	1.00	46.97	0
ATOM 3487 OG1 THR A 484 89.451 89.676 2.114 1.00 49.12	MOTA	3486	CB	THR	A	484	88.710	88.453	2.002	1.00	48.12	С
MICH 510, CCI IIII II COVICE COVICE COVICE	MOTA	3487	OG1	THR	A	484	89.451	89.676	2.114	1.00	49.12	0



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ATOM	3488	CG2	THR	A	484	87.219	88.767	1.861	1.00	47.18	С
ATOM	3489	N	ASN	Α	485	91.184	87.894	4.171	1.00	47.14	N
ATOM	3490	CA	ASN	Α	485	92.605	87.644	4.356	1.00	47.75	C
MOTA	3491	C	ASN	A	485	92.864	87.515	5.854	1.00	47.58	C
ATOM	3492	0	ASN	A	485	92.851	88.507	6.594	1.00	47.74	0
MOTA	3493	CB	ASN	A	485	93.436	88.785	3.767	1.00	48.75	C
MOTA	3494	CG	ASN	Α	485	94.934	88.540	3.890	1.00	50.65	C
MOTA	3495	OD1	ASN	Α	485	95.453	87.518	3.429	1.00	50.94	0
ATOM	3496	ND2	ASN	A	485	95.636	89.483	4.508	1.00	51.72	N
ATOM	3497	N	GLU			93.099	86.276	6.281	1.00	46.65	N
MOTA	3498	CA	GLU	Α	486	93.332	85.930	7.680	1.00	45.73	C
ATOM	3499	C	GLU			94.720	86.252	8.207	1.00	45.11	C
ATOM	3500	0	GLU			95.660	86.465	7.443	1.00	45.20	0
ATOM	3501	CB	GLU			93.057	84.441	7.867	1.00	46.18	C
ATOM	3502	CG	GLU			91.649	84.047	7.462		46.84	C
ATOM	3503	CD	GLU			91.503	82.556	7.259		47.48	C
ATOM	3504	OE1	GLU			90.380	82.105	6.946		47.47	0
ATOM	3505	OE2	GLU			92.518	81.838	7.402		47.64	0
ATOM	3506	N	ALA			94.835	86.284	9.530		44.77	N
ATOM	3507	CA	ALA			96.106	86.563	10.187		44.41	C
ATOM	3508	C	ALA			96.892	85.264	10.290			C
ATOM	3509	0	ALA			96.314	84.184	10.407		43.71	0
ATOM	3510	CB	ALA			95.867	87.143	11.583		44.17	C
ATOM	3511	N	PRO			98.227	85.350	10.227		45.67	N
ATOM	3512	CA	PRO	A	488	99.083	84.168	10.317		46.18	C
ATOM	3513	C			488	98.978	83.485	11.675		46.38	C
ATOM ATOM	3514 3515	O CB			488 488	99.572	82.434	11.892		47.49	0
ATOM	3516	CG	PRO	A		100.474 100.351	84.751 86.122	10.064 10.696		46.01 46.03	C
ATOM	3517	CD	PRO			99.059	86.551	10.034		46.59	C
ATOM	3517	N	ASN			98.218	84.075	12.587		46.12	N
ATOM	3519	CA	ASN			98.073	83.499	13.918		46.19	C
ATOM	3520	C	ASN			96.604	83.237	14.261		45.99	C
ATOM	3521	0	ASN			96.258	82.970	15.415		45.30	0
ATOM	3522	СВ	ASN			98.712	84.433	14.948		45.82	C
ATOM	3523	CG	ASN			97.953	85.736	15.101		47.05	C
ATOM	3524	OD1	ASN			97.335	86.229	14.152		47.80	0
ATOM	3525	ND2	ASN	Α	489	98.019	86.320	16.293		47.88	N
ATOM	3526	N	TYR	Α	490	95.741	83.327	13.253		45.80	N
ATOM	3527	CA	TYR	Α	490	94.319	83.069	13.447		45.86	С
ATOM	3528	С	TYR	Α	490	93.669	82.536	12.168		46.13	С
MOTA	3529	0	TYR	A	490	93.361	83.295	11.246	1.00	45.86	0
MOTA	3530	CB	TYR	Α	490	93.576	84.328	13.916	1.00	44.08	C
ATOM	3531	CG	TYR	A	490	92.148	84.013	14.302	1.00	42.82	С
ATOM	3532	CD1	TYR	A	490	91.871	83.222	15.419	1.00	41.48	C
ATOM	3533	CD2	TYR	A	490	91.080	84.402	13.489	1.00	42.13	C
MOTA	3534	CE1	TYR	A	490	90.570	82.816	15.709	1.00	41.83	C

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ATOM	3535	CE2	TYR	A	490	89.776	84.001	13.772	1.00	41.77	C
MOTA	3536	CZ	TYR	Α	490	89.529	83.204	14.881	1.00	41.35	C
MOTA	3537	OH	TYR	Α	490	88.248	82.771	15.141	1.00	40.05	0
MOTA	3538	N	SER	Α	491	93.475	81.220	12.135	1.00	47.86	N
MOTA	3539	CA	SER	Α	491	92.880	80.512	11.005	1.00	49.91	C
MOTA	3540	C	SER	A	491	93.583	80.739	9.661	1.00	51.25	C
MOTA	3541	0	SER	Α	491	94.408	81.674	9.551	1.00	53.12	0
MOTA	3542	CB	SER	A	491	91.402	80.881	10.887	1.00	50.03	C
MOTA	3543	OG	SER	Α	491	90.700	80.462	12.043	1.00	51.19	0
ATOM	3544	N	VAL	Α	492	93.288	79.975	8.714	1.00	52.00	N
TER	3545		VAL	Α	492						
HETATM	3546	P	IMP		500	96.855	67.484	18.145	1.00	36.20	P
HETATM	3547	01P	IMP		500	96.567	66.958	16.764	1.00	35.78	0
HETATM	3548	O2P	IMP		500	97.831	68.643	18.107	1.00	37.27	0
HETATM	3549	03 P	IMP		500	95.630	67.848	18.926	1.00	37.73	0
HETATM	3550	05*	IMP		500	97.576	66.348	19.098	1.00	39.55	0
HETATM	3551	C5*	IMP		500	96.906	65.102	19.379	1.00	43.52	C
HETATM	3552	C4 *	IMP		500	97.725	64.188	20.314	1.00	46.38	C
HETATM	3553	04*	IMP		500	96.849	63.056	20.510		47.93	0
HETATM	3554	C3*	IMP		500	99.128	63.615	19.896	1.00	47.80	C
		03*	IMP		500	100.283	64.165	20.566		48.27	0
HETATM		C2*	IMP		500	99.063	62.138	20.401		48.14	C
HETATM		02*	IMP		500	99.459	61.962	21.777		47.86	0
HETATM		C1*	IMP		500	97.568	61.820	20.391		48.84	C
HETATM		N9	IMP		500	97.172	60.952	19.187		49.35	N
HETATM		C8	IMP		500	97.769	60.762	17.934		49.94	C
HETATM		N7	IMP		500	96.960	59.811	17.195		49.47	N
HETATM		C5	IMP		500	95.903	59.489	18.092		49.72	C
HETATM		C6	IMP		500	94.774	58.548 57.868	17.851 16.852		49.49 49.17	0
HETATM		06 N1	IMP IMP		500 500	94.520 93.959	58.521	18.994		50.36	N
HETATM		N1 C2	IMP		500	94.155	59.256	20.190		50.39	C
HETATM HETATM		N3	IMP		500	95.182	60.067	20.130		49.12	N
HETATM		C4	IMP		500	96.016	60.150	19.259		49.43	C
HETATM		0	НОН		501	100.576	78.275	17.774		32.86	0
HETATM		0	НОН		503	91.095	64.490	36.206	1.00		0
HETATM		0	нон		504	94.498	58.629	33.377	1.00		o
HETATM		0	нон		505	121.994	54.382	32.283	1.00		0
HETATM		0	нон		506	103.325	52.600	39.850		30.92	0
HETATM		0	нон		508	114.082	79.200	14.020			0
HETATM		0	нон		509	111.782	75.357	9.474		41.74	0
HETATM		0	нон		510	130.431	69.550	6.520		33.07	0
HETATM		0	нон		511	139.704	72.872	5.502		28.29	0
HETATM		0	НОН		512	138.165	79.843	7.558		37.74	0
HETATM		0	нон		513	110.012	65.496	9.942		27.19	0
HETATM		0	нон		514	95.709	85.791	36.220	1.00	26.84	0
HETATM		0	нон		515	97.359	70.230	25.703	1.00	26.44	0

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HETATM	3582	0	нон	516	111.343	65.984	36.009	1.00 27.68	. 0
HETATM	3583	0	HOH	517	92.610	64.734	28.562	1.00 25.19	0
HETATM	3584	0	HOH	518	136.337	83.402	18.230	1.00 44.74	0
HETATM	3585	0	HOH	519	125.872	72.828	24.786	1.00 56.58	0
HETATM	3586	0	HOH	520	123.394	85.143	17.982	1.00 44.83	0
HETATM	3587	0	нон	521	114.051	75.607	10.822	1.00 28.33	0
HETATM	3588	0	нон	522	95.439	56.751	31.398	1.00 35.18	0
HETATM	3589	0	HOH	523	126.131	73.771	18.625	1.00 28.52	0
HETATM	3590	0	HOH	524	88.573	61.550	21.356	1.00 35.43	0
HETATM	3591	0	HOH	525	84.159	71.151	29.206	1.00 24.40	0
HETATM	3592	0	HOH	526	133.284	55.863	10.735	1.00 36.12	0
HETATM	3593	0	HOH	527	127.221	75.361	20.788	1.00 31.12	0
HETATM	3594	0	HOH	528	144.500	73.437	21.138	1.00 37.89	0
HETATM	3595	0	HOH	529	104.651	89.119	21.108	1.00 45.21	0
HETATM	3596	0	HOH	530	97.113	80.827	16.767	1.00 29.31	0
HETATM	3597	0	HOH	531	115.587	76.287	34.279	1.00 41.57	0
HETATM	3598	0	HOH	532	110.085	78.952	34.960	1.00 36.98	0
HETATM	3599	0	HOH	533	113.576	64.235	37.324	1.00 44.04	0
HETATM	3600	0	HOH	536	113.308	82.910	26.870	1.00 35.47	0
HETATM	3601	0	HOH	537	135.913	65.420	20.988	1.00 42.04	0
HETATM	3602	0	HOH	538	81.878	61.168	23.889	1.00 37.18	0
HETATM	3603	0	HOH	539	112.973	80.799	18.829	1.00 46.80	0
HETATM	3604	0	HOH	540	114.422	67.736	18.471	1.00 21.38	0
HETATM	3605	0	HOH	541	149.015	73.669	7.625	1.00 31.27	0
HETATM	3606	0	HOH	542	94.881	66.686	41.661	1.00 28.69	0
HETATM		0	нон	544	100.410	54.930	42.976	1.00 40.88	0
HETATM	3608	0	HOH	546	123.513	59.702	10.409	1.00 32.15	0
HETATM	3609	0	нон	547	146.434	72.633	24.076	1.00 36.35	0
HETATM		0	нон	548	88.886	71.737	36.970	1.00 30.08	0
HETATM		0	нон	549	107.710	87.920	18.571	1.00 46.77	0
HETATM		0	HOH	550	89.670	63.104	37.776	1.00 47.36	0
HETATM		0	нон	552	122.692	77.602	19.074	1.00 36.80	0
HETATM		0	НОН	553	98.993	52.226	5.651	1.00 57.87	0
HETATM		0	НОН	554	113.536	82.903	22.647	1.00 41.61	0
HETATM		0	нон	555	115.587	74.041	10.297	1.00 38.20	0
HETATM		0	нон	557	98.458	49.507	39.120	1.00 62.44	0
HETATM		0	НОН	558	112.138	63.863	8.254	1.00 24.96	0
HETATM		0	нон	560	118.847	70.191	28.617	1.00 45.38	0
HETATM		0	нон	561	146.469	68.284	24.114	1.00 37.59	0
HETATM		0	нон	562	128.262	63.501	17.485	1.00 38.06	0
HETATM		0	НОН	563	119.639	64.695	7.607	1.00 46.90	0
HETATM		0	НОН	564	103.490	70.881	36.574	1.00 31.68	
HETATM		0	HOH	565 567	98.314	86.799	37.005	1.00 30.78	0
HETATM		0	HOH	567 568	131.484	71.418	23.548 41.874	1.00 36.31 1.00 35.52	0
HETATM		0	HOH	568 569	76.343 126.027	59.075	12.269	1.00 33.32	0
HETATM		0	HOH	569 570	109.042	74.363	33.675	1.00 43.43	0
HETATM	3028	0	НОН	570	103.042	/4.303	33.073	T.00 J3.31	9

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HETATM	3629	0	нон	573	80.409	90.280	22.579	1.00	34.21		0
HETATM	3630	0	HOH	574	132.441	66.786	9.212	1.00	43.87	•	0
HETATM	3631	0	HOH	575	109.944	79.697	32.230	1.00	32.32	(	0
HETATM	3632	0	HOH	577	144.216	66.858	24.156	1.00	40.46	(	0
HETATM	3633	0	HOH	578	129.414	81.484	16.530	1.00	61.51	(	0
HETATM	3634	0	HOH	579	96.972	92.020	39.467	1.00	39.03		0
HETATM	3635	0	HOH	580	94.372	70.562	18.189	1.00	26.74	(	0
HETATM	3636	0	HOH	581	88.506	73.637	23.695	1.00	31.00	(	0
HETATM	3637	0	HOH	584	100.007	84.003	17.987	1.00	28.74	(	0
HETATM	3638	0	HOH	585	121.405	52.600	17.537	1.00	40.20	(	0
HETATM	3639	0	HOH	588	119.167	48.434	16.339	1.00	65.21	(	0
HETATM	3640	0	HOH	589	102.348	55.052	4.902	1.00	39.64	(	0
HETATM	3641	0	HOH	595	97.908	70.027	21.031	1.00	34.06		0
HETATM	3642	0	HOH	596	146.572	53.248	20.725	1.00	45.62	(	0
HETATM	3643	0	HOH	598	90.443	55.094	8.608	1.00	49.49		0
HETATM	3644	0	HOH	599	109.411	77.677	2.747	1.00	53.11		0
HETATM	3645	0	HOH	600	107.764	45.882	37.351	1.00	45.02	•	0
HETATM	3646	0	HOH	601	87.454	73.205	38.703	1.00	45.44	•	0
HETATM	3647	0	HOH	602	80.069	88.678	20.249	1.00	42.72		0
HETATM	3648	0	HOH	603	117.159	79.599	17.309	1.00	30.98		0
HETATM	3649	0	HOH	604	84.446	85.847	6.609	1.00	66.35		0
HETATM	3650	0	HOH	605	142.262	74.880	21.888	1.00	42.69		0
HETATM	3651	0	HOH	606	133.945	64.662	8.678	1.00	39.03	•	0
HETATM	3652	0	HOH	607	110.322	88.556	20.880	1.00	51.80	•	0
HETATM	3653	0	HOH	608	118.514	60.464	29.384	1.00	33.39	•	0
HETATM	3654	0	HOH	609	82.950	78.301	25.302	1.00	33.55	•	0
HETATM	3655	0	HOH	610	111.407	75.421	34.632	1.00	44.58	•	0
HETATM	3656	0	нон	612	96.558	63.438	23.644	1.00	34.09	(	0
HETATM	3657	0	HOH	613	122.627	63.063	23.597	1.00	26.90	(	0
HETATM	3658	0	HOH	614	131.169	69.077	14.358	1.00	34.02		0
HETATM	3659	0	HOH	618	96.690	54.179	40.860	1.00	32.89		0
HETATM	3660	0	HOH	619	126.711	73.763	15.905		32.41		0
HETATM	3661	0	нон	621	92.253	63.599	39.103		25.94		0
HETATM	3662	0	HOH	622	97.813	87.700	39.424		55.96		0
HETATM	3663	0	HOH	623	95.535	56.002	20.767		65.87		0
HETATM	3664	0	HOH	624	129.489	78.975	20.616		46.43		0
HETATM	3665	0	HOH	625	119.866	48.017	8.457	1.00	69.76		0
HETATM	3666	0	НОН	627	134.345	53.135	10.168		47.96		0
HETATM	3667	0	HOH	628	96.130	73.007	12.133		65.41		0
HETATM		0	HOH	629	87.800	55.819	15.251		45.94		0
HETATM	3669	0	HOH	630	112.422	70.233	37.715		55.93		0
HETATM		0	HOH	632	121.366	74.739	24.939		57.43		0
HETATM	3671	0	нон	633	101.115	80.540	12.424		53.67		0
HETATM		0	НОН	634	124.882	64.152	26.715		65.56		0
HETATM		0	НОН	635	97.480	87.029	5.748		41.53		0
HETATM		0	НОН	636	98.153	71.085	17.480		23.05		0
HETATM	3675	0	HOH	638	109.916	53.593	7.340	1.00	60.03	•	0

HETATM	3676	0	НОН	639	109.356	46.219	43.098	1.00	74.41	0
HETATM	3677	0	HOH	640	123.090	65.562	2.087	1.00	64.16	0
HETATM	3678	0	HOH	641	121.091	58.113	37.981	1.00	53.09	0
HETATM	3679	0	HOH	642	106.879	71.024	39.823	1.00	49.73	0
HETATM	3680	0	HOH	644	125.842	60.086	22.102	1.00	57.75	0
HETATM	3681	0	HOH	645	89.792	80.261	21.919	1.00	65.49	0
HETATM	3682	0	HOH	646	127.581	71.494	10.877	1.00	41.56	0
HETATM	3683	0	HOH	647	116.711	52.692	38.666	1.00	55.88	0
HETATM	3684	0	HOH	648	137.352	49.806	16.559	1.00	67.37	0
HETATM	3685	0	HOH	649	93.707	89.594	38.028	1.00	28.70	0
HETATM	3686	0	НОН	650	102.722	58.318	7.082	1.00	52.26	0
HETATM	3687	0	нон	651	99.494	73.756	-9.807	1.00	57.10	0
HETATM	3688	0	HOH	652	100.369	60.404	12.181	1.00	24.56	0
HETATM	3689	0	HOH	653	86.387	73.114	22.271	1.00	36.14	0
HETATM	3690	0	HOH	655	82.774	85.056	16.980	1.00	29.50	0
HETATM	3691	0	HOH	656	138.719	75.780	24.731	1.00	43.44	0
HETATM	3692	0	HOH	657	135.443	63.373	23.553	1.00	62.79	0
HETATM	3693	0	HOH	658	136.532	59.341	22.164	1.00	61.51	0
HETATM	3694	0	нон	659	112.228	46.754	14.166	1.00	44.56	0
HETATM	3695	0	нон	661	85.056	77.737	37.011	1.00	51.97	0
HETATM	3696	0	HOH	664	93.862	94.123	24.058	1.00	44.59	0
HETATM	3697	0	HOH	665	125.115	50.268	16.057	1.00	50.48	0
HETATM	3698	0	нон	666	109.792	82.218	39.304	1.00	41.41	0
HETATM	3699	0	нон	667	81.419	71.538	32.800	1.00	43.84	0
HETATM	3700	0	нон	671	100.872	91.979	23.613	1.00	64.05	0
HETATM	3701	0	НОН	672	121.924	64.730	9.693	1.00	40.99	0
HETATM	3702	0	HOH	673	103.164	53.450	45.794	1.00	46.76	0
HETATM	3703	0	нон	674	112.887	44.758	35.892	1.00	60.15	0
HETATM	3704	0	нон	675	121.226	52.298	40.410	1.00	59.96	0
HETATM	3705	0	HOH	676	114.778	79.883	12.588	1.00	53.72	0
HETATM	3706	0	HOH	677	111.493	44.375	26.336	1.00	44.91	0
HETATM	3707	0	HOH	680	125.672	77.196	7.641	1.00	62.63	0
HETATM	3708	0	HOH	681	149.427	68.734	21.594	1.00	69.45	0
HETATM	3709	0	HOH	682	130.498	68.890	11.409	1.00	43.07	0
HETATM	3710	0	HOH	684	97.027	74.301	-8.091	1.00	51.74	0
HETATM	3711	0	HOH	685	93.468	57.644	35.310	1.00	25.70	0
HETATM	3712	0	НОН	687	120.082	63.118	33.794	1.00	66.97	0
HETATM	3713	0	HOH	688	91.794	50.180	5.365	1.00	61.88	0
HETATM	3714	0	HOH	691	120.955	66.509	26.956	1.00	57.92	0
HETATM	3715	0	HOH	692	147.976	65.172	14.279	1.00	58.48	0
HETATM	3716	0	HOH	693	90.415	78.310	23.880	1.00	38.31	0
HETATM	3717	0	НОН	694	113.372	43.333	17.881	1.00	68.93	0
HETATM	3718	0	нон	695	101.223	90.113	38.606		45.49	0
HETATM	3719	0	нон	696	108.151	50.895	41.168		59.49	0
HETATM		0	НОН	697	90.431	44.244	14.620		43.55	0
HETATM		0	HOH	698	146.554	70.443	18.977		32.63	0
HETATM	3722	0	нон	702	107.324	89.479	37.117	1.00	65.35	0

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HETATM	3723	0	нон	706	151.406	55.047	15.937	1.00 5	56.70	C	)
HETATM	3724	0	HOH	707	101.778	67.697	-5.655	1.00 3	34.92	C	)
HETATM	3725	0	HOH	709	136.699	62.881	-10.241	1.00 5	3.86	C	)
HETATM	3726	0	HOH	710	115.523	70.698	9.393	1.00 3	35.01	C	)
HETATM	3727	0	HOH	714	140.987	80.163	24.272	1.00 6	55.37	C	)
HETATM	3728	0	HOH	715	144.845	70.181	8.359	1.00 4	15.96	C	)
HETATM	3729	0	HOH	716	127.420	64.712	10.814	1.00 5	50.93	C	)
HETATM	3730	0	HOH	717	112.548	85.955	35.733	1.00 6	53.37	C	)
HETATM	3731	0	нон	718	96.397	65.225	43.866	1.00 5	54.78	C	)
HETATM	3732	0	HOH	719	149.381	55.765	8.190		16.95	C	
HETATM	3733	0	HOH	723	115.502	77.990	9.376		16.76	C	
HETATM	3734	0	HOH	725	76.437	79.568	26.459		59.19	C	
HETATM		0	НОН	726	95.324	49.183	27.259		51.94	C	
HETATM		0	НОН	727	111.936	82.375	12.461		38.86	C	
HETATM		0	нон	728	133.312	81.928	11.453	1.00 5		C	
HETATM		0	нон	729	107.996	85.280	18.442	1.00 3		C	
HETATM		0	нон	730	148.848	63.651	-10.490		18.09	C	
HETATM		0	нон	733	134.306	63.018	10.766		31.65	C	
HETATM		0	нон	735	124.671	60.360	17.610		52.69	C	
HETATM		0	НОН	736	111.727	60.489	42.963		52.15	C	
HETATM		0	НОН	737	134.980	50.157	7.477		50.92	_	
HETATM		0	нон	738	146.654	76.277	6.833 19.814	1.00 4		C	
HETATM		0	НОН НОН	739 741	89.251 105.433	64.149 55.341	8.828	1.00 5		C	
HETATM HETATM		0	нон	749	88.458	78.199	19.817	1.00 6		C	
HETATM		0	нон	750	106.898	44.639	18.376	1.00 5		C	
HETATM		0	нон	751	105.309	68.078	49.132	1.00 6		C	
HETATM		0	нон	752	92.980	48.934	17.873	1.00 4		C	
HETATM		0	нон	753	100.420	53.758	15.446	1.00 6		C	
HETATM		0	нон	754	120.798	66.196	40.717	1.00 6		C	)
HETATM		0	НОН	755	108.406	89.679	12.448	1.00 €	51.48	C	)
HETATM		0	нон	757	132.463	72.528	-4.509	1.00 5	59.31	C	)
HETATM	3755	0	нон	761	127.038	77.545	21.661	1.00 4	17.91	C	)
HETATM	3756	0	HOH	762	106.459	50.413	17.617	1.00 5	59.90	C	)
HETATM	3757	0	HOH	765	119.622	72.534	26.691	1.00 5	51.38	C	)
HETATM	3758	0	HOH	766	115.174	72.450	5.241	1.00 6	58.42	C	)
HETATM	3759	0	HOH	768	105.322	87.067	41.471	1.00 6	58.21	C	)
HETATM	3760	0	HOH	770	105.218	41.445	29.836	1.00	53.03	C	)
HETATM	3761	0	HOH	771	83.989	78.004	39.580	1.00	57.38	C	
HETATM	3762	0	НОН	772	148.829	67.245	20.069	1.00 4		C	
HETATM	3763	0	HOH	773	106.544	84.270	11.229	1.00 4		C	
HETATM		0	НОН	775	105.699	39.006	23.992	1.00 6		C	
HETATM		0	нон	778	114.007	85.900	33.916	1.00 4		C	
HETATM		0	НОН	780	93.889	60.095	23.291	1.00			
HETATM		0	HOH	781 787	129.215	73.681	21.887	1.00		C	
HETATM		0	HOH	787 789	81.781	76.779	28.784	1.00 6			
HETATM	3/69	0	HOH	788	147.918	62.639	14.966	1.00		_	

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HETATM	3770	0	нон	789	79.084	63.731	35.792	1.00 60.46	0
HETATM	3771	0	HOH	791	106.162	72.361	37.935	1.00 51.27	0
HETATM	3772	0	HOH	793	113.985	72.209	7.836	1.00 54.33	0
HETATM	3773	0	HOH	794	147.561	84.941	11.984	1.00 59.95	0
HETATM	3774	0	HOH	796	117.356	86.976	18.333	1.00 57.65	0
HETATM	3775	0	HOH	797	96.398	58.891	24.526	1.00 48.25	0
HETATM	3776	0	HOH	799	113.947	81.311	24.929	1.00 48.99	0
HETATM	3777	0	HOH	800	130.861	56.862	14.557	1.00 67.81	0
HETATM	3778	0	нон	803	136.966	56.000	-10.559	1.00 66.93	0
HETATM	3779	0	нон	807	120.744	54.584	40.505	1.00 65.23	0
HETATM	3780	0	HOH	809	111.191	79.767	14.798	1.00 39.00	0
HETATM	3781	0	HOH	810	100.504	80.761	16.287	1.00 40.40	0
HETATM	3782	0	HOH	812	119.309	46.953	19.630	1.00 58.74	0
${\tt HETATM}$	3783	0	HOH	817	79.924	78.741	21.466	1.00 62.43	0
HETATM	3784	0	HOH	819	114.995	48.374	8.804	1.00 68.51	0
HETATM	3785	0	HOH	823	121.219	69.376	9.088	1.00 61.14	0
HETATM	3786	0	HOH	824	83.259	52.990	49.620	1.00 42.61	0
HETATM	3787	0	HOH	826	73.121	53.007	46.366	1.00 51.95	0
HETATM	3788	0	нон	829	125.131	57.072	24.888	1.00 43.39	0
HETATM	3789	0	HOH	830	117.478	81.142	25.399	1.00 63.97	0
HETATM	3790	0	HOH	832	118.060	80.843	9.442	1.00 68.90	0
HETATM	3791	0	нон	833	125.309	81.768	10.259	1.00 36.71	0
HETATM	3792	0	нон	834	106.669	69.383	1.728	1.00 62.38	0
HETATM	3793	0	HOH	836	134.415	57.557	20.513	1.00 51.26	0
HETATM	3794	0	HOH	837	138.774	48.063	14.052	1.00 62.75	0
HETATM	3795	0	HOH	838	105.034	90.698	36.793	1.00 43.22	0
HETATM	3796	0	HOH	839	94.179	62.599	22.253	1.00 39.73	0
HETATM		0	нон	840	102.012	56.721	46.229	1.00 49.59	0
HETATM		0	нон	842	129.445	55.023	4.305	1.00 44.96	0
HETATM		0	нон	843	95.363	50.937	5.969	1.00 60.45	0
HETATM	3800	0	нон	847	148.499	52.780	8.080	1.00 60.01	0
HETATM		0	НОН	852	90.466	70.795	38.792	1.00 60.50	0
HETATM		0	НОН	853	138.576	76.612	2.592	1.00 62.24	0
HETATM		0	НОН	855	116.588	74.760	7.223	1.00 60.72	0
HETATM		0	НОН	860	113.703	91.874		1.00 68.47	0
HETATM		0	нон	861	130.923	52.830	6.921	1.00 62.84	0
HETATM		0	нон	862	142.316	48.653	13.579	1.00 50.41	0
HETATM		0	нон	863	132.567	53.947	3.095	1.00 68.97	0
HETATM		0	НОН	865	100.473	47.680	26.060	1.00 65.28	0
HETATM		0	НОН	866	133.655	75.864	1.041 7.931	1.00 69.12 1.00 61.12	0
HETATM		0	нон	867	122.519	76.254			0
HETATM		0	HOH	868 869	84.905	74.842 52.884	20.730 0.797	1.00 46.62 1.00 66.79	0
HETATM		_	HOH	869 972	148.011	87.503	38.810	1.00 88.79	0
HETATM		0	HOH	872 973	94.647	92.083	26.520	1.00 58.25	0
HETATM		0	HOH	873 875	101.350 126.984	55.912	-0.651	1.00 62.62	0
HETATM		0	HOH	875 879		68.643		1.00 83.71	0
HETATM	2219	0	HOH	878	127.346	00.043	12.003	1.00 33.33	•

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HETATM	3817	0	нон	879	117.590	70.113	8.367	1.00	32.01	•	0
HETATM	3818	0	HOH	884	94.685	91.830	3.118	1.00	53.98	(	0
HETATM	3819	0	HOH	886	94.421	91.554	39.231	1.00	35.13	•	0
HETATM	3820	0	НОН	887	90.370	90.226	5.222	1.00	36.45	•	0
HETATM	3821	0	HOH	888	138.171	82.094	23.696		43.62		0
HETATM		0	HOH	890	145.344	74.873	18.144		52.45		0
HETATM	3823	0	HOH	891	86.699	56.553	44.193		59.80		0
HETATM		0	нон	898	110.253	51.388	39.073		64.37		0
HETATM	3825	0	нон	899	142.548	59.418	25.624		68.03		0
HETATM	3826	0	нон	902	96.309	63.463	47.551		68.27		0
HETATM	3827	0	НОН	904	103.052	43.719	26.788		64.56		0
HETATM		0	HOH	905	148.314	72.538	19.514		53.02		0 0
HETATM		0	HOH	906	115.081 111.660	80.764 74.882	15.768 5.430		37.40 36.15		0
HETATM		0	НОН НОН	907 908	91.410	88.940	39.058		40.21		0
HETATM HETATM		0	НОН	909	92.100	65.397	41.837		42.61		0
		0	НОН	910	135.015	70.210	22.164		32.10		0
HETATM	3834	0	нон	911	124.196	60.165	24.502		68.95		0
HETATM	3835	0	нон	912	104.972	48.595	38.399		50.34		0
HETATM	3836	Ö	нон	913	143.458	60.042	28.053		36.04		0
HETATM		0	нон	916	101.435	59.347	18.933		50.24		o
HETATM		0	нон	917	112.207	49.761	38.732		26.79		o
HETATM		0	нон	918	129.684	70.406	8.637		46.40	•	0
HETATM		0	нон	921	80.125	93.172	15.865		52.80	•	0
HETATM		0	нон	922	149.474	69.335	14.961		54.64	(	0
HETATM		0	нон	926	102.735	61.951	20.408	1.00	35.26	(	0
HETATM		0	нон	927	70.905	53.569	44.583	1.00	70.52	•	0
HETATM	3844	0	нон	929	114.913	79.874	33.698	1.00	57.49	•	0
HETATM	3845	0	нон	930	150.193	75.660	8.945	1.00	35.94	•	0
HETATM	3846	0	HOH	931	108.648	54.745	43.966	1.00	62.03	(	0
HETATM	3847	0	HOH	936	121.915	58.864	24.758	1.00	55.08	(	0
HETATM	3848	0	HOH	937	79.417	58.337	48.896	1.00	68.62	•	0
HETATM	3849	0	HOH	938	113.378	43.843	23.780	1.00	41.09	•	0
HETATM	3850	0	нон	939	120.219	59.743	27.131		44.37		0
HETATM	3851	0	HOH	941	105.234	64.396	-0.669		73.28		0
HETATM	3852	0	нон	942	141.057	80.868	-0.762		67.90		0
HETATM		0	НОН	944	119.413	75.184	6.755		98.67		0
HETATM		0	НОН	945	95.626	74.384	14.294		63.69		0
HETATM		0	нон	946	95.387	53.160	20.326		59.76		0
HETATM		0	нон	947	112.354	72.038	3.479		55.13		0
HETATM		0	НОН	961	116.417	63.126	34.645		53.04		0
HETATM		0	НОН	962	91.960	70.760	43.791		62.09		0
HETATM		0	HOH	964 966	80.831	84.466	14.233		66.83 43.56		0
HETATM		0	HOH	966 968	111.313 133.045	64.422 80.261	5.416 24.904		59.86		0
HETATM		0	нон нон	966 971	113.856	45.592	40.153		68.81		0
HETATM HETATM		0	НОН	973	149.882	57.333	6.661		61.17		0
ULTAIGU	2002	J	TOT	213	147.002	ددد.،د	0.001	1.00	UI.I/		_

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HETATM	3864	0	нон	974	114.479	45.337	33.893	1.00 48.82	2 0
HETATM	3865	0	нон	976	122.683	49.979	23.699	1.00 54.50	0
HETATM	3866	0	нон	977	107.449	93.037	22.262	1.00 67.52	2 0
HETATM	3867	0	HOH	980	71.623	55.665	43.963	1.00 50.43	3 0
HETATM	3868	0	HOH	983	81.115	79.690	16.340	1.00 68.34	Į O
HETATM	3869	0	HOH	984	146.095	68.641	21.063	1.00 40.33	0
HETATM	3870	0	HOH	989	84.591	87.656	11.809	1.00 61.62	2 0
HETATM	3871	0	нон	990	101.284	84.890	35.573	1.00 66.53	0
HETATM	3872	0	HOH	991	132.290	57.405	16.338	1.00 68.86	0
HETATM	3873	0	HOH	992	107.181	71.461	43.131	1.00 68.43	
HETATM	3874	0	HOH	996	121.732	77.353	22.459	1.00 56.10	
HETATM		0	нон	997	123.339	62.223	9.181	1.00 53.79	
HETATM		0	нон	999	118.564	57.129	2.150	1.00 64.38	
HETATM		0	нон	1002	113.406	85.261	25.792	1.00 54.44	
HETATM		0	нон	1003	132.676	51.930	17.206	1.00 68.65	
HETATM		0	нон	1006	82.100	75.518	31.280	1.00 51.37	
HETATM		0	нон	1007	91.217	86.172	10.703	1.00 68.50	
HETATM		0	HOH	1011	148.150	63.664	-4.949	1.00 66.39	
HETATM		0	нон	1012	108.584	47.618	13.690	1.00 63.35	
HETATM		0	HOH	1014	104.916	54.259	6.694	1.00 66.63	
HETATM		0	нон	1021	127.338	67.350	-0.507	1.00 68.53	
HETATM		0	нон	1024	100.255	43.755	35.224	1.00 49.55	
HETATM		0	HOH	1026	113.002	85.034	18.817	1.00 68.03	
HETATM		0	HOH	1027	74.446	56.955	41.184	1.00 34.26	
HETATM		0	нон	1032	123.923	66.490	27.749	1.00 51.93	
HETATM		0	нон	1037	105.661	94.018	14.310	1.00 48.05	
HETATM		0	нон	1045	85.110	67.600	42.845	1.00 61.60	
HETATM		0	нон	1049	72.485	57.802	45.989	1.00 68.63	
HETATM		0	нон	1051	104.785	74.784	39.154	1.00 60.69	
HETATM		0	нон	1053	104.639	40.347	34.518	1.00 61.33	
HETATM		0	нон	1054	142.840	80.523	20.021	1.00 68.20	
HETATM		0	нон	1056	123.658	55.426	39.072	1.00 68.03	
HETATM		0	нон	1057	122.409	54.809	6.777	1.00 68.72	
HETATM		0	НОН	1060	148.405	75.478	20.015	1.00 68.99	
HETATM		0	НОН	1066	101.285	46.434	21.329	1.00 69.66	
HETATM		0	НОН	1068	101.265	47.738	38.183	1.00 52.34	_
HETATM		0	НОН	1072	116.191	83.171 83.118	15.683 19.379	1.00 65.19	_
HETATM		0	HOH	1076	124.162 114.649	91.913	27.612	1.00 63.6	
HETATM		0	НОН	1077	131.138	72.022	1.639	1.00 65.80	
HETATM		0	HOH	1078		95.248	23.931	1.00 67.60	_
HETATM		0	НОН НОН	1079 1080	104.565 130.600	83.061	14.460	1.00 67.86	
HETATM		0	НОН	1080	108.024	57.385	44.494	1.00 68.4	_
HETATM HETATM		0	нон	1081	98.180	52.040	22.425	1.00 57.8	
HETATM		0	HOH	1095	123.035	48.662	12.121	1.00 37.3	
HETATM		0	НОН	1100	116.951	82.153	13.434	1.00 47.3	_
HETATM		0	НОН	1100	93.000	74.011	11.563		
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HETATM	3911	0	нон	1110	84.826	60.423	43.980	1.00	68.47	0
HETATM	3912	0	HOH	1114	95.995	48.302	8.024	1.00	46.89	0
HETATM	3913	0	HOH	1115	146.331	50.245	-5.119	1.00	58.03	0
HETATM	3914	0	НОН	1117	93.037	80.264	22.271	1.00	36.08	0
HETATM	3915	0	HOH	1118	127.120	59.841	19.240	1.00	36.51	0
HETATM	3916	0	нон	1123	130.326	80.122	18.283	1.00	54.14	0
HETATM	3917	0	нон	1127	122.797	71.467	7.444		61.33	0
HETATM		0	нон	1128	88.326	63.775	40.835		65.83	0
HETATM		0	нон	1129	129.934	60.674	26.265		67.06	0
HETATM		0	нон	1132	91.126	55.593	11.580		55.91	0
HETATM		0	нон	1136	131.604	57.222	-9.584		62.55	0
HETATM		0	нон	1137	122.963	68.163	16.106	1.00	33.32	0
HETATM		0	нон	1140	107.890	88.486	9.887		61.56	0
HETATM		0	нон	1142	143.969	58.642	-10.289		69.08	0
HETATM		0	нон	1143	88.527	78.564	-1.195	1.00		0
HETATM		0	HOH	1146	109.850	50.588	43.199		64.27	0
HETATM		0	HOH	1151	112.701 102.584	76.952 50.394	6.277 12.484	1.00	55.30 59.53	0
HETATM		0	нон нон	1154 1161	95.856	79.770	13.615	1.00		0
HETATM HETATM		0	НОН	1161	149.220	72.694	15.463	1.00	36.11	0
HETATM		0	HOH	1162	134.026	86.608	28.831	1.00		0
HETATM		0	НОН	1168	137.288	47.676	-0.383		58.53	0
HETATM		0	НОН	1169	96.461	76.369	-1.039		40.84	0
HETATM		0	нон	1170	146.839	76.924	18.226		62.40	Ō
HETATM		0	нон	1173	84.778	62.413	46.009		56.40	0
HETATM		0	нон	1174	104.665	61.328	-0.147		68.18	0
HETATM		0	нон	1176	148.238	49.557	15.253		68.84	0
HETATM		0	нон	1180	96.826	57.686	5.466		68.82	0
HETATM		0	нон	1181	97.848	45.596	18.230	1.00	68.53	0
HETATM		0	нон	1183	105.561	78.152	46.280	1.00	55.13	0
HETATM		0	нон	1184	148.363	55.663	18.453	1.00	67.78	0
HETATM	3942	0	нон	1188	117.761	72.763	3.201	1.00	49.39	0
HETATM	3943	0	нон	1190	129.206	55.861	-5.442	1.00	69.86	0
HETATM	3944	0	нон	1195	107.481	76.284	39.087	1.00	57.19	0
HETATM	3945	0	нон	1206	122.685	66.549	4.934	1.00	62.97	0
HETATM	3946	0	HOH	1207	150.879	41.687	5.427	1.00	67.80	0
HETATM	3947	0	HOH	1216	134.077	45.934	8.123	1.00	68.54	0
HETATM	3948	0	нон	1217	92.702	54.498	4.335	1.00	68.29	0
HETATM	3949	0	HOH	1227	133.307	89.606	16.031	1.00	55.60	0
HETATM	3950	0	HOH	1228	145.314	58.907	23.524		57.48	0
HETATM	3951	0	нон	1231	121.333	47.473	28.343		42.57	0
HETATM	3952	0	нон	1237	80.672	64.102	43.307		62.67	0
HETATM	3953	0	НОН	1239	79.197	76.085	29.202		42.02	0
HETATM		0	НОН	1240	147.532	79.940	6.435		49.20	0
HETATM		0	НОН	1241	105.341	73.622	43.925		46.80	0
HETATM		0	НОН	1242	108.748	84.315	11.317		43.50	0
HETATM	3957	0	HOH	1243	113.748	76.205	13.481	1.00	31.05	0



HETATM	3958	0	нон	1244	106.486	82.249	41.211	1.00	53.73	(	0
HETATM	3959	0	HOH	1245	123.229	67.990	30.863	1.00	67.62	(	0
HETATM	3960	0	HOH	1246	97.244	56.293	3.245	1.00	59.53		0
HETATM	3961	0	нон	1247	84.115	75.748	18.158		47.44		0
HETATM		0	HOH	1248	92.641	62.480	43.494		56.54		0
HEŤATM	3963	0	HOH	1249	126.850	67.707	7.524		63.22		0
HETATM	3964	0	HOH	1250	116.737	46.525	9.414		60.31		0
HETATM		0	нон	1251	99.435	55.524	20.442		68.76		0
HETATM		0	нон	1252	93.533	48.432	11.284		64.31		0
HETATM		0	НОН	1253	115.458	55.820	8.527		68.91		0
HETATM		0	нон	1254	94.383	48.132	30.166		55.54		0
HETATM		0	нон	1255	136.004	53.964	17.602		50.55		0
HETATM		0	нон	1256	97.765	60.337	0.278		67.66		0
HETATM		0	НОН	1257	81.887	70.128	40.015		60.06		0
HETATM		0	НОН	1258	98.568	43.853	36.969		60.96		0
HETATM		0	НОН	1259	102.312	50.226	23.207		69.15 62.17		0
HETATM		0	нон нон	1260 1261	93.845 122.247	73.542 50.835	7.463 30.996		66.32		0
HETATM HETATM		0	нон нон	1261	137.839	46.740	1.638		44.22		0
HETATM		0	нон	1263	107.295	79.492	3.520		56.86		0
HETATM		0	нон	1264	107.233	49.640	21.504		48.82		0
HETATM		0	нон	1265	105.132	63.518	48.797		62.13		0
HETATM		0	нон	1266	139.420	62.113	23.787		50.94		0
HETATM		o	нон	1267	144.043	77.286	3.516		68.14		0
HETATM		0	НОН	1268	149.733	53.900	4.381		56.17		0
HETATM		0	НОН	1269	103.004	91.675	22.454		51.28	(	0
HETATM		0	нон	1270	102.342	79.977	8.282	1.00	60.86	(	0
HETATM	3985	0	нон	1271	104.432	79.198	8.137	1.00	49.45		0
HETATM	3986	0	нон	1272	96.642	78.325	15.154	1.00	49.69	(	0
HETATM	3987	0	HOH	1273	123.113	83.532	10.129	1.00	46.69	(	0
HETATM	3988	0	нон	1274	108.924	75.712	35.457	1.00	44.24	(	0
HETATM	3989	0	HOH	1275	120.284	52.133	13.839	1.00	49.42	(	0
HETATM	3990	0	HOH	1276	153.804	67.675	3.008	1.00	68.53	(	0
HETATM	3991	0	HOH	1277	132.756	49.791	2.618		54.77		0
HETATM	3992	0	нон	1278	123.687	61.097	-1.686		66.47		0
HETATM	3993	0	НОН	1279	79.098	85.995	16.502		68.34		0
HETATM		0	HOH	1280	81.604	77.273	22.663		51.85		0
HETATM		0	нон	1281	97.665	46.523	31.377		66.98		0
HETATM		0	нон	1282	124.226	53.393	42.604		56.65		0
HETATM		0	нон	1283	70.053	51.433	45.374		31.76		0
HETATM		0	нон	1284	133.004	59.283	22.298		51.14		0 0
HETATM		0	НОН	1285	110.435	58.745	9.141		66.24		0
HETATM		0	HOH	1286	131.690	83.699	10.424		50.89		0
HETATM		0	HOH	1287	87.121	83.954 70.654	6.897 47.251		61.10 59.87		0
HETATM HETATM		0	нон нон	1289 1290	103.343 151.878	57.545	47.251		68.93		0
HETATM		0	НОН	1290	109.757	52.533	5.140		68.61		0
DETAIM	<b>3004</b>	9	поп	1691	109.757	J4.JJJ	2.140	1.00	00.01	· ·	_



HETATM	4005	0	HOH	1292	137.500	85.244	14.713	1.00 48.53	0
HETATM	4006	0	HOH	1293	99.481	43.592	30.277	1.00 56.76	0
HETATM	4007	0	HOH	1294	79.393	66.499	44.205	1.00 51.03	0
HETATM	4008	0	HOH	1295	93.025	76.731	12.952	1.00 68.06	0
HETATM	4009	0	HOH	1296	104.177	39.836	37.064	1.00 66.28	0
HETATM	4010	0	HOH	1297	131.482	71.092	26.769	1.00 63.53	0
HETATM	4011	0	HOH	1299	108.732	64.733	46.862	1.00 59.61	0
HETATM	4012	0	HOH	1300	85.693	84.234	8.773	1.00 61.29	0
HETATM	4013	0	HOH	1301	130.439	55.137	19.928	1.00 68.52	0
HETATM	4014	0	HOH	1302	126.942	81.225	22.497	1.00 68.12	0
HETATM	4015	0	HOH	1303	85.867	53.208	45.199	1.00 54.32	0
HETATM	4016	0	HOH	1304	104.487	89.634	40.115	1.00 64.91	0
HETATM	4017	0	HOH	1305	106.217	68.163	-1.625	1.00 55.54	0
HETATM	4018	0	HOH	1306	105.015	51.028	40.376	1.00 45.33	0
HETATM	4019	0	HOH	1307	120.170	70.835	39.052	1.00 58.73	0
HETATM	4020	0	HOH	1308	121.326	61.115	30.539	1.00 67.23	0
HETATM	4021	0	нон	1309	107.923	60.643	46.488	1.00 69.06	0
HETATM	4022	0	HOH	1310	78.786	53.486	48.325	1.00 55.45	0
HETATM	4023	0	HOH	1311	130.804	55.401	22.465	1.00 59.73	0
HETATM	4024	0	HOH	1312	150.487	50.345	12.892	1.00 61.66	0
HETATM		0	HOH	1313	100.834	38.834	32.531	1.00 62.67	0
HETATM	4026	0	нон	1314	111.244	65.102	1.801	1.00 68.69	0
HETATM	4027	0	HOH	1315	132.117	79.062	22.385	1.00 40.28	0
HETATM	4028	0	нон	1316	91.048	57.371	42.981	1.00 57.35	0
HETATM	4029	0	HOH	1317	144.712	49.327	0.134	1.00 49.54	0
HETATM	4030	0	HOH	1318	147.789	61.584	-9.156	1.00 68.51	0
HETATM		0	нон	1319	123.090	61.674	6.437	1.00 69.05	0
HETATM		0	НОН	1320	116.358	60.623	35.690	1.00 69.30	0
HETATM		0	нон	1321	113.530	91.177	21.954	1.00 52.62	0
HETATM		0	нон	1322	110.177	74.769	14.221	1.00 65.83	0
HETATM		0	нон	1323	135.219	50.902	-6.112	1.00 68.00	0
HETATM		0	нон	1324	121.026	52.096	9.382	1.00 68.04	0
HETATM		0	нон	1325	86.410	84.214	10.939	1.00 54.69	0
HETATM		0	нон	1326	92.066	62.207	-0.340	1.00 62.69	0
HETATM		0	нон	1327	108.159	72.272	1.047	1.00 69.41	0
HETATM		0	нон	1328	136.575	47.899		1.00 67.46	. 0
HETATM		0	НОН	1329	112.693	55.745	39.945	1.00 50.31	0
HETATM		0	HOH	1330	142.437		-12.250	1.00 63.99	0
HETATM		0	HOH	1331	97.845	53.121	3.695	1.00 68.33	0
HETATM		0	HOH	1332	135.048	60.985	20.232	1.00 43.95 1.00 35.53	0
HETATM		0	HOH	1334	131.683	75.485	22.258	1.00 53.53	0
HETATM		0.	HOH	1335	105.140	83.991	8.520 31.673	1.00 42.24	0
HETATM		0	НОН НОН	1336 1337	119.421 100.568	70.763 48.021	23.330	1.00 42.24	0
HETATM		0				54.355		1.00 49.89	. 0
HETATM HETATM		0	нон нон	1338 1339		47.835		1.00 63.64	. 0
			НОН	1340		62.937			o
HETATM	# ADT	0	HOH	T240	120.201	02.937	13.900	1.00 05.00	J

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1.00 47.23

1.00 53.04

1.00 57.31

1.00 48.23

1.00 44.60

1.00 59.34

1.00 52.82

1.00 68.35

1.00 48.04

1.00 50.57

1.00 44.69

8.757

3.434

7.299

3.914

7.477

7.356

13.378

33.425

24.673

16.877

49.080

32.787

32.009

15.005

27.038

44.276

### -125-

51.862

62.277

79.489

48.324

79.259

54.202

83.848

76.586

97.127

77.928

89.092

64.700

72.862

90.948

75.697

95.254

152.210

111.310

149.427

111.003

116.893

117.820

111.270

129.000

124.726

93.181

79.571

115.741

85.780

112.933

86.047

92.864

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HETATM	4055	0	HOH	1344
HETATM	4056	0	HOH	1345
HETATM	4057	0	HOH	1346
HETATM	4058	0	нон	1347
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HETATM	4060	0	HOH	1349
HETATM	4061	0	HOH	1350
HETATM	4062	0	HOH	1351
HETATM	4063	0	HOH	1352
HETATM	4064	0	HOH	1353
HETATM	4065	0	HOH	1354
HETATM	4066	0	HOH	1356
MTATM	4067	0	HOH	1357
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CONECT	404	403	405	407
CONECT	405	404	406	
CONECT	406	405		
CONECT	407	404	408	
CONECT	408	407	409	
CONECT	409	408	410	
CONECT	410	409		
CONECT	459	460		
CONECT	460	459	461	463
CONECT	461	460	462	
CONECT	462	461		
CONECT	463	460	464	
CONECT	464	463	465	
CONECT	465	464	466	
CONECT	466	465		
CONECT	576	577		
CONECT	577	576	578	580
CONECT	578	577	579	
CONECT	579	578		
CONECT	580	577	581	
CONECT	581	580	582	
CONECT	582	581	583	
CONECT	583	582		
CONECT	882	883		
CONECT	883	882		886
CONECT	884	883	885	
CONECT	885	884	_	
CONECT	886	883	887	
CONECT	887	886		
CONECT	888	887	889	

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TABLE 7

889 888

CONECT 1096 1095 1097

CONECT 1095 1094 1096 1098

CONECT 1094 1095

CONECT

#### CONECT 1097 1096 CONECT 1098 1095 1099 CONECT 1099 1098 1100 CONECT 1100 1099 1101 CONECT 1101 1100 CONECT 1211 1212 CONECT 1212 1211 1213 1215 CONECT 1213 1212 1214 CONECT 1214 1213 CONECT 1215 1212 1216 CONECT 1216 1215 1217 CONECT 1217 1216 1218 CONECT 1218 1217 CONECT 2707 2708 CONECT 2708 2707 2709 2711 CONECT 2709 2708 2710 CONECT 2710 2709 CONECT 2711 2708 2712 CONECT 2712 2711 2713 CONECT 2713 2712 2714 CONECT 2714 2713 CONECT 2733 2734 CONECT 2734 2733 2735 2737 CONECT 2735 2734 2736 CONECT 2736 2735 CONECT 2737 2734 2738 CONECT 2738 2737 2739 CONECT 2739 2738 2740 CONECT 2740 2739 CONECT 2934 2935 CONECT 2935 2934 2936 2938 CONECT 2936 2935 2937 CONECT 2937 2936 CONECT 2938 2935 2939 CONECT 2939 2938 2940

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CONECT 2974 2971 2975

CONECT 2975 2974 2976

CONECT 2971 2970 2972 2974

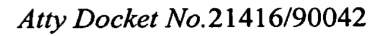
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CONECT 2970 2971

CONECT 2973 2972



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CONECT 2977 2976
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CONECT 3160 3159 3161
CONECT 3161 3160
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CONECT 3163 3162 3164
CONECT 3164 3163 3165
CONECT 3165 3164
CONECT 3211 3212
CONECT 3212 3211 3213 3215
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CONECT 3214 3213
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CONECT 3216 3215 3217
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CONECT 3546 3547 3548 3549 3550
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CONECT 3550 3546 3551
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CONECT 3553 3552 3558
CONECT 3554 3552 3555 3556
CONECT 3555 3554
CONECT 3556 3554 3557 3558
CONECT 3557 3556
CONECT 3558 3553 3556 3559
CONECT 3559 3558 3560 3568
CONECT 3560 3559 3561
CONECT 3561 3560 3562
CONECT 3562 3561 3563 3568
CONECT 3563 3562 3564 3565
CONECT 3564 3563
CONECT 3565 3563 3566
CONECT 3566 3565 3567
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TABLE 7

CONECT 3567 3566 3568

CONECT 3568 3559 3562 3567

MASTER 437 0 14 18 16 0 1 6 4066 1 127 37

END

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15

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35

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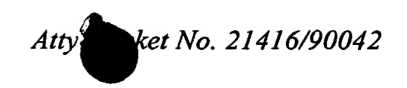
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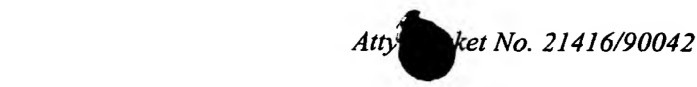
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